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(57) Abstract

The present invention is concerned with the identification of genes or functional fragments thereof from Candida albicans which are critical for growth and cell division and which genes may be used as selective drug targets to treat Candida albicans associated infections. Novel nucleic acid sequences from Candida albicans are also provided and which encode the polypeptides which are critical for growth of Candida albicans. Methods for the identification of anti-fungal compounds which inhibit fungal or yeast growth are also contemplated.

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DRUG TARGETS IN CANDIDA ALBICANS

The present invention is concerned with the identification of genes or functional fragments thereof from Candida albicans which are critical for growth and cell division and which genes may be used as selective drug targets to treat Candida albicans associated infections. Novel nucleic acid sequences from Candida albicans are also provided and which encode the polypeptides which are critical for growth of Candida albicans.

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Opportunistic infections in immunocompromised hosts represent an increasingly common cause of mortality and morbidity. Candida species are among the most commonly identified fungal pathogens associated with such opportunistic infections, with Candida albicans being the most common species. Such fungal infections are thus problematical in, for example, AIDS populations in addition to normal healthy women where Candida albicans yeasts represent the most common cause of vulvovaginitis.

Although compounds do exist for treating such disorders, such as, amphotericin, these drugs are generally limited in their treatment because of their toxicity and side effects. Therefore, there exists a need for new compounds which may be used to treat Candida associated infections in addition to compounds which are selective in their action against Candida albicans.

Classical approaches for identifying anti-fungal compounds have relied almost exclusively on inhibition of fungal or yeast growth as an endpoint. Libraries of natural products, semi-synthetic, or synthetic chemicals are screened for their ability to kill or arrest growth of the target pathogen or a related nonpathogenic model organism. These tests are

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cumbersome and provide no information about a compound's mechanism of action. The promising lead compounds that emerge from such screens must then be tested for possible host-toxicity and detailed mechanism of action studies must subsequently be conducted to identify the affected molecular target.

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The present inventors have now identified a range of nucleic acid sequences from Candida albicans which encode polypeptides which are critical for its survival and growth. These sequences represent novel targets which can be incorporated into an assay to selectively identify compounds capable of inhibiting expression of such polypeptides and their potential use in alleviating diseases or conditions associated with Candida albicans infection.

Therefore, according to a first aspect of the invention there is provided a nucleic acid molecule encoding a polypeptide which is critical for survival and growth of the yeast *Candida albicans* and which nucleic acid molecule comprises any of the sequences of nucleotides illustrated in any of Sequence ID Nos. 1 to 9.

whilst the molecules defined herein have been established as being critical for growth and metabolism of Candida albicans, for some of the molecules no apparent functionality has been assigned by virtue of the fact that no functionally related sequences in other prokaryctic or eukaryotic organism can be found in respective databases. Thus, advantageously these sequences may be species specific in which case they may be used be used as selective targets for treatment of diseases mediated by Candida Albicans infection. Thus, in one aspect of the invention the nucleic acid molecules preferably comprise the sequences identified in sequence ID Nos. 1, 4, 5 to 9.

In another aspect of the invention the sequences have been arranged functionally and of nucleotides illustrated in Sequence ID Nos. 2 or 3 are preferred and even more preferably in Sequence ID No. 2 and fragments or derivatives of said nucleic acid molecules.

Letters utilised in the sequences according to the invention which are not recognisable as letters of the genetic code signify a position in the nucleic acid sequence where one or more of bases A, G, C or T can occupy the nucleotide position. Representative letters used to identify the range of bases which can be used are as follows:

15	М:	A or C
	R:	A or G
	W :	A or T
	S:	C or G
	Y:	C or T
20	K:	G or T
	v:	A or C or G
	Н:	A or C or T
		A or G or T
	В:	C or G or T
25	N:	G or A or T or C

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In one embodiment of each of the above identified aspects of the invention the nucleic acid may comprise a mRNA molecule or alternatively a DNA and preferably a cDNA molecule.

Also provided by the present invention is a nucleic acid molecule capable of hybridising to the nucleic acid molecules illustrated in any of Figures 1 to 9 under high stringency conditions such as antisense molecule and which conditions are generally known to those of skill in the art.

Stringency of hybridisation as used herein refers

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to conditions under which polynucleic acids are stable. The stability of hybrids is reflected in the melting temperature (Tm) of the hybrids. Tm can be approximated by the formula:

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81.5°C+16.6(Log₁₀[Na⁺]+0.41 (%G&C)-600L/L

wherein L is the length of the hybrids in nucleotides. Tm decreases approximately by 1-1.5°C with every 1% decrease in sequence homology.

The term "stringency" refers to the hybridisation conditions wherein a single-stranded nucleic acid joins with a complementary strand when the purine or pyramidine bases therein pair with their corresponding base by hydrogen bonding. High stringency conditions favour homologous base pairing whereas low stringency conditions disfavour non-homologous base pairing.

"Low stringency" conditions comprise, for example, a temperature of about 37°C or less, a formamide concentration of less than about 50%, and a moderate to low salt (SSC) concentration; or, alternatively, a temperature of about 50°C or less, and a moderate to high salt (SSPE) concentration, for example 1M NaCl.

"High stringency" conditions comprise, for example, a temperature of about 42°C or less, a formamide concentration of less than about 20%, and a low salt (SSC) concentration; or, alternatively, a temperature of about 65°C, or less, and a low salt (SSPE) concentration. For example, high stringency conditions comprise hybridization in 0.5 M NaHPC4, 7% sodium dodecyl sulfate (SDS), 1 mM EDTA at 65°C (Ausubel, F.M. et al. Current Protocols in Molecular Biology, Vol. I, 1989; Green Inc. New York, at 2.10.3).

"SSC" comprises a hybridization and wash solution. A stock 20% SSC solution contains 3M sodium

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chloride, 0.3M sodium citrate, pH 7.0.

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"SSPE" comprises a hybridization and wash solution. A 1X SSPE solution contains 180 mM NaCl, 9mM Na₂HPO₄ and 1 mM EDTA, pH 7.4.

The nucleic acid capable of hybridising to nucleic acid molecules according to the invention will generally be at least 70%, preferably at least 80 or 90% and more preferably at least 95% homologous to the nucleotide sequences illustrated in any of Figures 1 to 9.

The DNA molecules according to the invention may, advantageously, be included in a suitable expression vector to express polypeptides encoded therefrom in a suitable host which are critical for growth and survival of Candida albicans.

An expression vector according to the invention includes a vector having a nucleic acid according to the invention operably linked to regulatory sequences, such as promoter regions, that are capable of effecting expression of said DNA fragments. The term "operably linked" refers to a juxtaposition wherein the components described are in a relationship permitting them to function in their intended manner. Such vectors may be transformed into a suitable host cell to provide for expression of a polypeptide according to the invention. Thus, in a further aspect, the invention provides a process for preparing polypeptides according to the invention which comprises cultivating a nost cell, transformed, transfected or infected with an expression vector as described above under conditions to provide for expression by the vector of a coding sequence encoding the polypeptides, and recovering the expressed polypeptides.

The vectors may be, for example, plasmid, virus or phage vectors provided with an origin of replication, optionally a promoter for the expression.

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of said nucleotide and optionally a regulator of the promoter. The vectors may contain one or more selectable markers, such as, for example, ampicillin resistance.

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Regulatory elements required for expression include promoter sequences to bind RNA polymerase and transcription initiation sequences for ribosome binding. For example, a bacterial expression vector may include a promoter such as the lac promoter and for translation initiation the Shine-Dalgarno sequence and the start codon AUG. Similarly, a eukaryotic expression vector may include a heterologous or homologous promoter for RNA polymerase II, a downstream polyadenylation signal, the start codon AUG, and a termination codon for detachment of the ribosome. Such vectors may be obtained commercially or assembled from the sequences described by methods well known in the art.

Polynucleotides according to the invention may be inserted into the vectors described in an antisense orientation in order to provide for the production of antisense RNA. Antisense RNA or other antisense nucleic acids may be produced by synthetic means.

In accordance with the present invention, a defined nucleic acid includes not only the identical nucleic acid but also any minor base variations including in particular, substitutions in bases which result in a synonymous codon (a different codon specifying the same amino acid residue) due to the degenerate code. The term "nucleic acid sequence" also includes the complementary sequence to any single stranded sequence given regarding base variations.

The present invention also comprises within its scope proteins or polypeptides expressed by the nucleic acid molecules according to the invention or a functional equivalent, derivative or bioprecursor thereof.

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The present invention also advantageously provides nucleic acid sequences of at least approximately 10 contiguous nucleotides of a nucleic acid according to the invention and preferably from 10 to approximately 120 nucleotides. In another aspect of the invention, nucleotide acid sequences are provided from 10 to 50 nucleotides. These sequences may, advantageously be used as probes or primers to initiate replication, or the like. Such nucleic acid sequences may be produced according to techniques well known in the art, such as by recombinant or synthetic means. They may also be used in diagnostic kits or the like for detecting the presence of a nucleic acid according to the invention. These tests generally comprise contacting the probe with the sample under hybridising conditions and detecting for the presence of any duplex or triplex formation between the probe and any nucleic acid in the sample.

According to the present invention, these probes may be anchored to a solid support. Preferably, they are present on an array so that multiple probes can simultaneously hybridize to a single biological sample. The probes can be spotted onto the array or synthesized in situ on the array. See Lockhart et al., Nature Biotechnology, Vol. 14, December 1996, "Expression monitoring by hybridization to high-density oligonucleotide arrays." A single array can contain more than up to more than a million different probes in discrete locations.

Advantageously, the nucleic acid sequences, according to the invention may be produced using such recombinant or synthetic means, such as for example using PCR cloning mechanisms which generally involve making a pair of primers, which may be between approximately 10 to 120 nucleotides to a region of the gene which is desired to be cloned, bringing the primers into contact with mRNA, cDNA, or genomic DNA

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from a cell, performing a polymerase chain reaction under conditions which bring about amplification of the desired region, isolated the amplified region or fragment and recovering the amplified DNA. Generally, such techniques as defined herein are well known in the art, such as described in Sambrook et al (Molecular Cloning: a Laboratory Manual, 1989).

The nucleic acids or oligonucleotides according to the invention may carry a revealing label.

Suitable labels include radioisotopes such as ³²P or ³⁹S, enzyme labels or other protein labels such as biotin or fluorescent markers. Such labels may be added to the nucleic acids or oligonucleotides of the invention and may be detected using known techniques per se.

The polypeptide or protein according to the invention includes all possible amino acid variants encoded by the nucleic acid molecule according to the invention including a polypeptide encoded by said molecule and having conservative amino acid changes. Polypeptides according to the invention further include variants of such sequences, including naturally occurring allelic variants which are substantially homologous to said polypeptides. In this context, substantial homology is regarded as a sequence which has at least 70%, preferably 80 or 90% amino acid homology with the polypeptides encoded by the nucleic acid molecules according to the invention.

Nucleic acids and polypeptides which are particularly preferred are those comprising the sequences of nucleotides illustrated in figures 1 to 3 and polypeptides illustrated in figures 14 to 16. However, a particularly preferred nucleic acid comprises the sequences of nucleotides illustrated in Figures 2 and/or 3, and their corresponding amino acid sequences identified in Figures 15 and 16.

Nucleotide sequences according to the invention

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are particularly advantageous as selective therapeutic targets for treating Candida albicans associated infections. For example, an antisense nucleic acid capable of binding to the nucleic acid sequence illustrated in any of Figures 1 to 9 may be used to selectively inhibit expression of the corresponding polypeptides, leading to impaired growth of the Candida albicans with reductions of associated illnesses or diseases.

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The nucleic acid molecule or the polypeptide according to the invention may be used as a medicament, or in the preparation of a medicament, for treating diseases or conditions associated with Candida albicans infection.

Advantageously, the nucleic acid molecule or the polypeptide according to the invention may be provided in a pharmaceutical composition together with a pharmaceutically acceptable carrier, diluent or excipient therefor.

The present invention is further directed to inhibiting expression of nucleic acids according to the invention in vivo by the use of antisense technology. Antisense technology can be used to control gene expression through triple-helix formation of antisense DNA or RNA, both of which methods are based on binding of a polynucleotide to DNA or RNA. For example, the 5' coding portion or the mature protein sequence, which encodes for the protein of the present invention, is used to design an antisense RNA oligonuclectide of from 10 to 50 base pairs in length. A DNA oliganucleotide is designed to be complementary to a region of the gene involved in transcription (triple-helix - see Lee et al. Nucl. Acids Res., 6:3073 (1979); Cooney et al., Science, 241:456 (1988); and Dervan et al., Science, 251: 1360 (1991), thereby preventing transcription and the production of the

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corresponding protein. The antisense RNA oligonucleotide hybridises to the mRNA in vivo and blocks translation of an mRNA molecule into the corresponding protein (antisense - Okano, J. Neurochem., 56:560 (1991); Oligodeoxynucleotides as Antisense Inhibitors of Gene Expression, CRC Press, Boca Raton, FL (1988)).

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Antibodies to the protein or polypeptide of the present invention may, advantageously, be prepared by techniques which are known in the art. For example, polyclonal antibodies may be prepared by inoculating a host animal, such as a mouse, with the polypeptide according to the invention or an epitope thereof and recovering immune serum. Monoclonal antibodies may be prepared according to known techniques such as described by Kohler R. and Milstein C., Nature (1975) 256, 495-497.

Antibodies according to the invention may also be used in a method of detecting for the presence of a polypeptide according to the invention, which method comprises reacting the antibody with a sample and identifying any protein bound to said antibody. A kit may also be provided for performing said method which comprises an antibody according to the invention and means for reacting the antibody with said sample.

Proteins which interact with the polypeptide of the invention may be identified by investigating protein-protein interactions using the two-hybrid vector system first proposed by Chien et al. (1991).

This technique is based on functional reconstitution in vivo of a transcription factor which activates a reporter gene. More particularly the technique comprises providing an appropriate host cell with a DNA construct comprising a reporter gene under the control of a promoter regulated by a transcription factor having a DNA binding domain and an activating domain, expressing in the host cell a first hybrid DNA

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sequence encoding a first fusion of a fragment or all of a nucleic acid sequence according to the invention and either said DNA binding domain or said activating domain of the transcription factor, expressing in the host at least one second hybrid DNA sequence, such as a library or the like, encoding putative binding proteins to be investigated together with the DNA binding or activating domain of the transcription factor which is not incorporated in the first fusion; detecting any binding of the proteins to be investigated with a protein according to the invention by detecting for the presence of any reporter gene product in the host cell; optionally isolating second hybrid DNA sequences encoding the binding protein.

An example of such a technique utilises the GAL4 protein in yeast. GAL4 is a transcriptional activator of galactose metabolism in yeast and has a separate domain for binding to activators upstream of the galactose metabolising genes as well as a protein binding domain. Nucleotide vectors may be constructed, one of which comprises the nucleotide residues encoding the DNA binding domain of GAL4. These binding domain residues may be fused to a known protein encoding sequence, such as for example the nucleic acids according to the invention. vector comprises the residues encoding the protein binding domain of GAL4. These residues are fused to residues encoding a test protein. Any interaction between polypeptides encoded by the nucleic acid according to the invention and the protein to be tested leads to transcriptional activation of a reporter molecule in a GAL4 transcription deficient yeast cell into which the vectors have been transformed. Preferably, a reporter molecule such as B-galactosidase is activated upon restoration of transcription of the yeast galactose metabolism genes. Further provided by the present invention is one

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or more *Candida albicans* cells comprising an induced mutation in the DNA sequence encoding the polypeptide according to the invention.

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A further aspect of the invention provides a method of identifying compounds which selectively inhibit or interfere with the expression, the functionality of polypeptides expressed from the nucleotides sequences illustrated in any of Figures 1 to 9 or the metabolic pathways in which these polypeptides are involved and which are critical for growth and survival of Candida albicans, which method comprises (a) contacting a compound to be tested with one or more Candida albicans cells having a mutation in a nucleic acid molecule according to the invention which mutation results in overexpression or underexpression of said polypeptides in addition to one or more wild type Candida cells, (b) monitoring the growth and/or activity of said mutated cell compared to said wild type wherein differential growth or activity of said one or more mutated Candida cells provides an indication of selective action of said compound on said polypeptide or another polypeptide in the same or a parallel pathway.

Compounds identifiable or identified using the method according to the invention, may advantageously be used as a medicament, or in the preparation of a medicament to treat diseases or conditions associated with Candida albicans infection. These compounds may also advantageously be included in a pharmaceutical composition together with a pharmaceutically acceptable carrier, diluent or excipient therefor.

A further aspect of the invention provides a method of identifying DNA sequences from a cell or organism which DNA encodes polypeptides which are critical for growth or survival, which method comprises (a) preparing a cDNA or genomic library from

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said cell or organism in a suitable expression vector which vector is such that it can either integrate into the genome in said cell or that it permits transcription of antisense RNA from the nucleotide sequences in said cDNA or genomic library, (b) selecting transformants exhibiting impaired growth and determining the nucleotide sequence of the cDNA or genomic sequence from the library included in the vector from said transformant. Preferably, the cell or organism may be any yeast or filamentous fungus, such as, for example, Saccharomyces cerevisiae, Schizosaccharomyces pombe or Candida albicans.

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A further aspect of the invention provides a pharmaceutical composition comprising any of a compound, an antisense molecule or an antibody according to the invention together with a pharmaceutically acceptable carrier, diluent or excipient therefor.

The antisense molecules or indeed the compounds identified as agonists or antagonists of the nucleic acids or polypeptides according to the invention may be used in the form of a pharmaceutical composition, which may be prepared according to procedures well known in the art. Preferred compositions include a pharmaceutically acceptable vehicle or diluent or excipient, such as for example, a physiological saline solution. Other pharmaceutically acceptable carriers including other non-toxic salts, sterile water or the like may also be used. A suitable buffer may also be present allowing the compositions to be lyophilized and stored in sterile conditions prior to reconstitution by the addition of sterile water for subsequent administration. Incorporation of the polypeptides of the invention into a solid or semisolid biologically compatible matrix may be carried out which can be implanted into tissues requiring treatment.

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The carrier can also contain other pharmaceutically acceptable excipients for modifying other conditions such as pH, osmolarity, viscosity, sterility, lipophilicity, solubility or the like. Pharmaceutically acceptable excipients which permit sustained or delayed release following administration may also be included.

The polypeptides, the nucleic acid molecules or compounds according to the invention may be administered orally. In this embodiment they may be encapsulated and combined with suitable carriers in solid dosage forms which would be well known to those skilled in the art.

As would be well known to those of skill in the art, the specific dosage regime may be calculated according to the body surface area of the patient or the volume of body space to be occupied, dependent upon the particular route of administration to be used. The amount of the composition actually administered will, however, be determined by a medical practitioner, based on the circumstances pertaining to the disorder to be treated, such as the severity of the symptoms, the composition to be administered, the age, weight, and response of the individual patient and the chosen route of administration.

The present invention may be more clearly understood with reference to the accompanying example, which is purely exemplary, with reference to the accompanying drawings, wherein

Figures 1 and 2:

are nucleotide sequences isolated from Candida albicans and which have an identified function based on sequence homology with proteins from other organisms and which sequences are not present in the public domain.

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5	Figures 3 :	is a nucleotide sequence isolated from Candida albicans and which has an identified function based on sequence homology with proteins from other organisms and which sequence is partially present in the public domain.
10	Figures 4 :	is a nucleotide sequence of previously unknown function isolated from <i>Candida albicans</i> and which is partially present in the public domain.
15	Figures 5 to 9:	are nucleotide sequences of previously unknown function isolated from Candida albicans.
20	Figure 10 :	is a diagrammatic representation of plasmid pGAL1PNiST-1.
	Figure 11	is a nucleotide sequence of plasmid pGAL1PNiST-1 of Figure 10.
25	Figure 12 :	is a diagrammatic representation of plasmid pGAL1PSiST-1.
30 35	Figure 13 :	is a nucleotide sequence of plasmid pGAL1PSiST-1 of Figure 12.
	Figures 14 to 20:	are amino acid sequences of the appropriately corresponding DNA sequences illustrated in Figures 1 to 9 with reference to Table 1.
	Figures 21 to 27:	are growth curves of Candida albicans strains showing antisense

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induced reduction in growth.

Figures 28 to 31: are growth curves of Candida albicans strains including knock-outs in the relevant gene identified.

Example 1

Identification of novel drug targets in C. albicans by anti-sense and disruptive integration

The principle of the approach is based on the fact that when a particular *C. albicans* mRNA is inhibited by producing the complementary anti-sense RNA, the corresponding protein will decrease. If this protein is critical for growth or survival, the cell producing the anti-sense RNA will grow more slowly or will die.

Since anti-sense inhibition occurs at mRNA level, the gene copy number is irrelevant, thus allowing applications of the strategy even in diploid organisms.

Anti-sense RNA is endogenously produced from an integrative or episomal plasmid with an inducible promoter; induction of the promoter leads to the production of an RNA encoded by the insert of the plasmid. This insert will differ from one plasmid to another in the library. The inserts will be derived from genomic DNA fragments or from cDNA to cover-to the extent possible- the entire genome.

The vector is a proprietary vector allowing integration by homologous recombination at either the homologous insert or promoter sequence in the Candida genome. After introducing plasmids from cDNA or genomic libraries into C. albicans, transformants are screened for impaired growth after promoter (& thus anti-sense) induction in the presence of lithium

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acetate. Lithium acetate prolongs the G1 phase and thus allows anti-sense to act during a prolonged period of time during the cell cycle. Transformants which show impaired growth in both induced and non-induced media, thus showing a growth defect due to integrative disruption, are selected as well.

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Transformants showing impaired growth are supposed to contain plasmids which produce anti-sense RNA to mRNAs critical for growth or survival. Growth is monitored by measuring growth-curves over a period of time in a device (Bioscreen Analyzer, Labsystems) which allows simultaneous measurement of growth-curves of 200 transformants.

Subsequently plasmids can be recovered from the transformants and the sequence of their inserts determined, thus revealing which mRNA they inhibit. In order to be able to recover the genomic or cDNA insert which has integrated into the Candida genome, genomic DNA is isolated, cut with an enzyme which cuts only once into the library vector (and estimated approx. every 4096 bp in the genome) and religated. PCR with primers flanking the insert will yield (partial) genomic or cDNA inserts as PCR fragments which can directly be sequenced. This PCR analysis (on ligation reaction) will also show us how many integrations occurred. Alternatively the ligation reaction is transformed to E. coli and PCR analysis is performed on colonies or on plasmid DNA derived thereof.

This method is employed for a genome-wide search for novel *C. albicans* genes which are important for growth or survival.

Materials & Methods

Construction of pGallPNiST-1

The backbone of the pGAL1PNiST-1 vector (integrative anti-sense Sfil-Notl vector) is

pGEM11Zf(+) (Promega Inc.). First, the CaMAL2

EcoRI/SalI promoter fragment from pDBV50 (D.H. Brown et al. 1996) was ligated into EcoRI/SalI-opened pGEM11Zf(+) resulting in the intermediate construct pGEMMAL2P-1. Into the latter (MscI/CIP) the CaURA3 selection marker was cloned as a Eco47III/XmnI fragment derived from pRM2. The resulting pGEMMAL2P-2 vector was NotI/HindIII opened in order to accept the NotI-stuffer-SfiI cassette from pPCK1NiSCYCT-1 (EagI/HindIII fragment): pMAL2PNiST-1. Finally, the plasmid pGAL1PNiST-1 was constructed by exchanging the SalI/Ecl136II MAL2 promoter in pMAL2PNiST-1 by the XhoI/Smal GAL1 promoter fragment derived from pRM2GAL1P.

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Construction of pGallPSiST-1

The vector pGAL1PSiST-1 was created for cloning the small genomic DNA fragments (flanked by SfiI sites) behind the GALl promoter. The only difference with pGAL1PNiST-1 is that the hIFNß (stuffer fragment) 20 insert fragment in pGAL1PSiST-1 is flanked by two SfiI sites in stead of a Sfil and a Notl site as in pGAL1PNiST-1. To construct pGAL1PSiST-1 the EcoRI-HindIII fragment, containing hIFN\$ flanked by a SfiI and a NotI site, of pMAL2pHiET-3 (unpublished) was 25 exchanged by the EcoRI-HindIII fragment, containing hIFN β flanked by two SfiI sites, from YCp50S-S (an E. coli / S. cerevisiae shuttle vector derived from the plasmid YCp50, which is deposited in the ATCC collection (number 37419; Thrash et al., 1985); an 30 EcoRI-HindIII fragment, containing the gene hIFNβ, which is flanked by two SfiI sites, was inserted in YCp50, creating YCp50S-S), resulting into plasmid pMAL2PSiST-1. The MAL2 promoter from pMAL2PSiST-1 (by a Nael-ball digest) was further replaced by the GAL1 35 promoter from pGAL1PNi3T-1 (via a XhoI-FSPI digest),

creating the vector pGAL1PSiST-1.

Candida albicans genomic library

* Preparation of the genomic DNA fragments A Candida albicans genomic DNA library with small DNA 5 fragments (400 to 1,000 bp) was prepared. Genomic DNA of Candida albicans B2630 was isolated following a modified protocol of Blin and Stafford (1976). The quality of the isolated genomic DNA was checked by gel electrophoresis. Undigested DNA was located on the gel 10 above the marker band of 26,282 bp. A little smear, caused by fragmentation of the DNA, was present. To obtain enrichment for genomic DNA fragments of the desired size, the genomic DNA was partially digested. Several restriction enzymes (AluI, HaeIII and RsaI; 15 all creating blunt ends) were tried out. The appropriate digest conditions have been determined by titration of the enzyme. Enrichment of small DNA fragments was obtained with 70 units of AluI on 10 μg of genomic DNA for 20 min. T4 DNA polymerase 20 (Boehringer) and dNTPs (Boehringer) were added to polish the DNA ends. After extraction with phenolchloroform the digest was size-fractionated on an agarose gel. The genomic DNA fragments with a length of 500 to 1,250 bp were eluted from the gel by 25 centrifugal filtration (Zhu et al., 1985). SfiI adaptors (5' GTTGGCCTTTT) or (5' AGGCCAAC) were attached to the DNA ends (blunt) to facilitate cloning of the fragments into the vector. Therefore, a 8-mer and 11-mer oligonucleotide (comprising the SfiI site) 30 were kinatesed and annealed. After ligation of these adaptors to the DNA fragments a second sizefractionation was performed on an agarose gel. DNA fragments of 400 to 1150 bp were eluted from the gel by centrifugal filtration. 35

* Preparation of the pGAL1PSiST-1 vector fragment

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The small genomic DNA fragments were cloned after the GAL1 promoter in the vector pGAL1PSIST-1. Qiagen-purified pGAL1PSiST-1 plasmid DNA was digested with SfiI and the largest vector fragment eluted from the gel by centrifugal filtration (Zhu et al., 1985). Ligation with a control DNA fragment, flanked by SfiI sites, was performed as a control. The ligation mix was electroporated to MC1061 E. coli cells. Plasmid DNA of 24 clones was analyzed. In all cases the control fragment was inserted in the pGAL1PSiST-1 vector fragment.

* Upscaling

All genomic DNA fragments (450 ng) were ligated into the pGAL1PSiST-1 vector (20 ng). After electroporation at 2500V, 40µF circa 400,000 clones were obtained. These clones were pooled into three groups and stored as glycerol slants. Also Qiagen-purified DNA was prepared from these clones. A clone analysis showed an average insert length of 600 bp and a percentage of 91 for clones with an insert. The size of the library corresponds to 5 times the diploid genome. The genomic DNA inserts are sense or antisense orientated in the vector.

25 Candida albicans cDNA library

Total RNA was extracted from Candida albicans
B2630 grown on respectively minimal (SD) and rich
(YPD) medium as described by Chirgwin et al. in
Sambrook et al 1996. mRNA was prepared from total RNA
using the Invitrogen Fast Track procedure.

First strand cDNA is synthesised with the Superscript Reverse Transcriptase (BRL) and with an oligo dT-NotI Primer adapter. After second strand synthesis, cDNA is polished with Klenow enzyme and purified over a Sephacryl S-400 spun column. Phosphorylated SfiI adapters are then ligated to the

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cDNA, followed by digestion with the *Not*I restriction enzyme. The *SfiI/Not*I cDNA is then purified and sized on a Biogel column A150M.

First fraction contains approximately 38,720 clones by transformation, the second fraction only 1540 clones. Clone analysis:
Fr. I: 22/24 inserts, 16 3 1000 bp, 4 3 2000 bp, average size: 1500 bp.
Fr. II: 9/12 inserts, 3 3 1000 bp, average size: 960 bp cDNA was ligated in a NotI/SfiI opened pGAL1PNiST-1

Candida transformation

vector (anti-sense)

The host strain used for transformation is a C.

albicans ura3 mutant, CAI-4, which contains a deletion in orotidine-5'-phosphate decarboxylase and was obtained from William Fonzi, Georgetown University (Fonzi and Irwin). CAI-4 was transformed with the above described cDNA library or genomic library using the Pichia spheroplast module (Invitrogen). Resulting transformants were plated on minimal medium supplemented with glucose (SD, 0.67% or 1.34% Yeast Nitrogen base w/o amino acids + 2% glucose) plates and incubated for 2-3 days at 30°C.

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Screening for mutants

Starter cultures were set up by inoculating each colony in 1 ml SD medium and incubating overnight at 30°C and 300 rpm. Cell densities were determined using a Coulter counter (Coulter Z1; Coulter electronics limited). 250.000 cells/ml were inoculated in 1 ml SD medium and cultures were incubated for 24 hours at 30°C and 300 rpm. Cultures were washed in minimal medium without glucose (S) and the pellet resuspended in 650 μ l S medium. 8 μ l of this culture is used for inoculating 400 μ l cultures in a Honeywell-100 plate

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(Bioscreen analyzer; Labsystems). Each transformant was grown during three days in S medium containing LiAc; pH 6.0, with 2% glucose/2% maltose or 2% galactose/2% maltose respectively while shaking every 3 minutes for 20 seconds. Optical densities were measured every hour during three consecutive days and growth curves were generated (Bioscreen analyzer; Labsystems).

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Growth curves of transformants grown in respectively anti-sense non-inducing (glucose/maltose) and inducing (galactose/maltose) medium are compared and those transformants showing impaired growth upon anti-sense induction are selected for further analysis. Transformants showing impaired growth by virtue of integration into a critical gene are also selected.

Isolation of genomic or cDNA inserts

Putatively interesting transformants are grown in 1.5 ml SD overnight and genomic DNA is isolated using the Nucleon MI Yeast kit (Clontech). Concentration of genomic DNA is estimated by analyzing a sample on an agarose gel.

20 ng of genomic DNA is digested for three hours with an enzyme that cuts uniquely in the library vector (SacI for the genomic library; PstI for the cDNA library) and treated with RNAse. Samples are phenol/chloroform extracted and precipitated using NaOAc/ethanol.

The resulting pellet is resuspended in 500 μ l ligation mixture (1 x ligation buffer and 4 units of T4 DNA ligase; both from Boehringer) and incubated overnight at 16°C.

After denaturation (20 min 65°C), purification (phenol/chloroform extraction) and precipitation (NaOAc/ethanol) the pellet is resuspended in 10 μ l MilliQ (Millipore) water.

PCR analysis

Inverse PCR is performed on 1 μ l of the precipitated ligation reaction using library vector specific primers (oligo23 5' TGC-AGC-TCG-ACC-TCG-ACT-G 3' and oligo25 5' GCG-TGA-ATG-TAA-GCG-TGA-C 3' for the 5 genomic library; 3pGALNistPCR primer :5'TGAGCAGCTCGCCGTCGCGC 3' and 5pGALNistPCR primer: 5'GAGTTATACCCTGCAGCTCGAC 3' for the cDNA library; both from Eurogentec) for 30 cycles each consisting of (a) 1 min at 95 °C, (b) 1 min at 57 °C, and (c) 3 min at 10 72 °C. In the reaction mixture 2.5 units of Taq polymerase (Boehringer) with TaqStart antibody (Clontech) (1:1) were used, and the final concentrations were 0.2 $\mu \mathrm{M}$ of each primer, 3 mM MgCl2 (Perkin Elmer Cetus) and 200 μM dNTPs (Perkin Elmer 15 Cetus). PCR was performed in a Robocycler (Stratagene).

Sequence determination

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20 Resulting/PCR products were purified using PCR purification kit (Qiagen) and were quantified by comparison of band intensity on EtBr stained agarose gel with the intensity of DNA marker bands. The amount of PCR product (expressed in ng) used in the sequencing reaction is calculated as the length of the PCR product in basepairs divided by 10. Sequencing reactions were performed using the ABI Prism BigDye Terminator Cycle Sequencing Ready Reaction Kit according to the instructions of the manufacturer (PE Applied Biosystems, Foster City, CA) except for the following modifications.

The total reaction volume was reduced to 15 μ l. Reaction volume of individual reagents were changed accordingly. 6.0 μ l Terminator Ready Reaction Mix was replaced by a mixture of 3.0 μ l Terminator Ready Reaction Mix + 3.0 μ l Half Term (GENPAK Limited,

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Brighton, UK). After cycle sequencing, reaction mixtures were purified over Sephadex G50 columns prepared on Multiscreen HV opaque microtitex-plates (Millipore, Molsheim, Fr) and were dried in a speedVac. Reaction products were resuspended in 3 μl loading buffer. Following denaturation for 2 min at 95°C, 1 μ l of sample was applied on a 5% Long Ranger Gel (36 cm well-to-read) prepared from Singel Packs according to the supplier's instructions (FMC BioProducts, Rockland, ME). Samples were run for 7 hours 2X run on a ABI 377XL DNA sequencer. collection version 2.0 and Sequence analysis version 3.0 (for basecalling) software packages are from PE Applied Biosystems. Resulting sequence text files were copied onto a server for further analysis. 15

Sequence analysis

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Nucleotide sequences were imported in the VectorNTI software package (InforMax Inc, North Bethesda, MD, USA), and the vector and insert regions 20 of the sequences were identified. Sequence similarity searches against public and commercial sequence databases were performed with the BLAST software package (Altschul et al., 1990) version 1.4. Both the original nucleotide sequence and the six-frame 25 conceptual translations of the insert region were used as query sequences. The used public databases were the EMBL nucleotide sequence database (Stoesser et al., 1998), the SWISS-PROT protein sequence database and its supplement TrEMBL (Bairoch and Apweiler, 1998), 30 and the ALCES Candida albicans sequence database (Stanford University, University of Minnesota). The commercial sequence databases used were the LifeSeq® human and PathoSeqÔ microbial genomic databases (Incyte Pharmaceuticals Inc., Palo Alto, CA, USA), and 35 the GENESEQ patent sequence database (Derwent, London,

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UK). Three major results were obtained on the basis of the sequence similarity searches: function, novelty, and specificity. A putative function was deduced on the basis of the similarity with sequences with a known function, the novelty was based on the absence or presence of the sequences in public databases, and the specificity was based on the similarity with vertebrate homologues.

10 Methods

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Blastx of the nucleic acid sequences against the appropriate protein databases: Swiss-Prot for clones of which the complete sequence is present in the public domain, and paorfp (PathoSeq $^{\text{TM}}$) for clones of which the complete sequences is not present in the public domain.

The protein to which the translated nucleic acid sequence corresponds to is used as a starting point. The differences between this protein and our translated nucleic acid sequences are marked with a double line and annotated above the protein sequence. The following symbols are used:

a one-letter amino acid code or the ambiguity code X is used if our translated nucleic acid sequence has another amino acid on a certain position,

the stop codon sign * is used if our translated nucleic acid sequence has a stop codon on a certain position,

The letters fs (frame shift) are used if a frame shift occurs in our translated nucleic acid sequence, and another reading frame is used,

the words ambiguity or ambiguities are used if a part of our translated nucleic acid sequence is present in the proteins, but not visible in the alignments of the blast results,

The phrase "missing sequence" is used if the translated nucleic acid sequence does not comprise

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that part of the protein.

Blastx: compares the six-frame conceptual translation products of a nucleotide query sequence (both strands) against a protein sequence database.

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Gene Knock-outs

To verify that the growth effect was due to the interference with the identified gene and to support the specificity of the antisense effect, single allele knock-outs were made in the identified genes (Figures 28 to 31) using the URA-blaster method (Fonzi and Irwin 1993).

Screening for compounds modulating expression of polypeptides critical for growth and survival of C. albicans

The method proposed is based on observations (Sandbaken et al., 1990; Hinnebusch and Liebman 1991; Ribogene PCT WO 95/11969, 1995) suggesting that underexpression or overexpression of any component of a process (e.g. translation) could lead to altered sensitivity to an inhibitor of a relevant step in that process. Such an inhibitor should be more potent against a cell limited by a deficiency in the macromolecule catalyzing that step and/or less potent against a cell containing an excess of that macromolecule, as compared to the wild type (WT) cell.

Mutant yeast strains, for example, have shown that some steps of translation are sensitive to the stoichiometry of macromolecules involved. (Sandbaken et al. 1996). Such strains are more sensitive to compounds which specifically perturb translation (by acting on a component that participates in translation) but are equally sensitive to compounds with other mechanisms of action.

This method thus not only provides a means to

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identify whether a test compound perturbs a certain process but also an indication of the site at which it exerts its effect. The component which is present in altered form or amount in a cell whose growth is affected by a test compound is potentially the site of action of the test compound.

The assay to be set up involves measurement of growth of an isogenic strain which has been modified only in a certain specific allele, relative to a wild type (WT) *C. albicans* strain, in the presence of R-compounds. Strains can be ones in which the expression of a specific essential protein is impaired upon induction of anti-sense or strains which carry disruptions in an essential gene. An in silico approach to finding novel essential genes in *C. albicans* will be performed. A number of essential genes identified in this way will be disrupted (in one allele) and the resulting strains can be used for comparative growth screening.

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Assay for High Throughput screening for drugs

35 μ l minimal medium (S medium + 2% galactose + 2% maltose) is transferred in a transparent flat-bottomed 96 well plate using an automated pipetting system (Multidrop, Labsystems). A 96-channel pipettor (Hydra, Robbins Scientific) transfers 2.5 μ l of R-compound at 10⁻³ M in DMSO from a stock plate into the assay plate.

The selected C. albicans strains (mutant and parent (CAI-4) strain) are stored as glycerol stocks (15%) at -70°C. The strains are streaked out on selective plates (SD medium) and incubated for two days at 30°C. For the parent strain, CAI-4, the medium is always supplemented with 20 μ g/ml uridine. A single colony is scooped up and resuspended in 1 ml minimal medium (S medium + 2% galactose + 2% maltose). Cells

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are incubated at 30°C for 8 hours while shaking at 250 rpm. A 10 ml culture is inoculated at 250.000 cells/ml. Cultures are incubated at 30°C for 24 hours while shaking at 250 rpm. Cells are counted in Coulter counter and the final culture (S medium + 2% galactose + 2% maltose) is inoculated at 20.000 to 50.000 cells/ml. Cultures are grown at 30°C while shaking at 250 rpm until a final OD of 0.24 (+/- 0.04) 600nM is reached.

200 μ l of this yeast suspension is added to all wells of MW96 plates containing R-compounds in a 450 (or 250) μ l total volume. MW96 plates are incubated (static) at 30°C for 48 hours.

Optical densities are measured after 48 hours.

Test growth is expressed as a percentage of positive control growth for both mutant (x) and wild type (y) strains. The ratio (x/y) of these derived variables is calculated.

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Table 1

	Seq ID No.	Figure No.	Clone '*-	Function
	1	1	382c_cp	-
5	2	2	392c_cp	TUF1
	3	3	-	RAD53
	4	4	417c_cpG2L	-
	5	5	323c_af	-
	6	6	322c_cp1	-
10	7	7	26g3	-
	8	8	409c_cp	_
	9	9	382c_cpG1L2	-
	10	14	382c_cp (prt)	-
	11	15	392c_cp (prt)	TUF1
15	12	16		RAD53
	13	17	325c_af (prt) ²	_
	14	18	322c_cp (prt) ²	-
	15	19	26g3 (prt)	-
	16	20	417c_cp 92L (prt)	-

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1. 322c-cp is a member of the UPF0057 protein family. It contains potential transmembrane regions (6-23aa; 30-53aa) and could be low temperature or salt-stress inducible.

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2. 325c-af shows similarity to IMP4 yeast and related proteins and it might be involved in rRNA processing in Candida albicans in a similar way to IMP4.

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Claims

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- 1. A nucleic acid molecule encoding a polypeptide which is critical for survival and growth of the yeast *Candida albicans* and which nucleic acid molecule comprises any of the sequences of nucleotides illustrated in Seq ID Nos 1 to 9.
- 2. A nucleic acid molecule encoding a polypeptide which is critical for survival and growth of the yeast *Candida albicans* and which nucleic acid molecule comprises any of the sequences of nucleotides illustrated in Seq ID Nos 1 to 3.
- 15 3. A nucleic acid molecule encoding a polypeptide which is critical for survival and growth of the yeast *Candida albicans* and which nucleic acid molecule comprises any of the sequences of nucleotides illustrated in Seq ID Nos 1 or 2 and fragments or derivatives of said nucleic acid molecules.
 - 4. A nucleic acid molecule according to any of claims 1 to 3 which is mRNA.
- 5. A nucleic acid molecule according to any of claims 1 to 3 which is DNA.
 - 6. A nucleic acid molecule according to claim 5 which is cDNA.
 - 7. A nucleic acid molecule capable of hybridising to the molecules according to any of claims 1 to 6 or the sequences illustrated in any of Seq ID Nos 1 to 9 under high stringency conditions.
 - 8. An antisense molecule comprising a nucleic

acid molecule capable of hybridising to the molecules according to any of claims 1 to 6 or the sequences illustrated in any of Seq ID Nos 1 to 9.

- 9. Cells containing a nucleic acid molecule according to any of claims 1 to 8, wherein said cells are bacterial or eukaryotic.
- 10. A polypeptide encoded by the nucleic acid 10 molecule according to any of claims 1 to 7 or the sequences illustrated in any of Seq ID Nos 1 to 9.
 - 11. A polypetide having any of amino acid sequences illustrated in any of Seq ID Nos 14 to 20.
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 12. A recombinant DNA construct comprising a nucleic acid molecule according to claim 5 or 6.
- 13. A recombinant DNA construct comprising a nucleic acid molecule according to claim 5 or 6 wherein said nucleic acid molecule is inserted in the antisense crientation.
- 14. A recombinant DNA construct according to 25 claim 12 or 13 wherein said recombinant DNA construct is an expression vector.
 - 15. A construct according to claim 14 which comprises an inducible promoter.

- 16. A construct according to claim 14 or 15 which comprises a sequence encoding a reporter molecule.
- 35 17. Cells containing a recombinant DNA construct according to any of claims 12 to 16, wherein said cells are bacterial or eukaryotic.

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18. A nucleic acid molecule according to any of claims 1 to 8 or the nucleotide sequences illustrated in Seq ID Nos 1 to 9 for use as a medicament.

- 19. Use of a nucleic acid molecule according to any of claims 1 to 8 or the sequences illustrated in Seq ID Nos 1 to 9 in the preparation of a medicament for treating Candida albicans associated diseases.
- 10 20. A polypeptide according to claim 10 or 11 for use as a medicament.
- 21. Use of a polypeptide according to claim 10 or 11 in the preparation of a medicament for treating Candida albicans associated infections.
 - 22. A pharmaceutical composition comprising a nucleic acid molecule according to any of claims 1 to 8 or a polypeptide according to claim 10 or 11 together with a pharmaceutically acceptable carrier diluent or excipient therefor.

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- 23. A Candida albicans cell comprising an induced mutation in the DNA sequence encoding the polypeptide according to claim 10.
- 24. A method of identifying compounds which selectively modulate expression or functionality of polypeptides or metabolic pathways in which these polypeptides are involved and which are crucial for growth and survival of *Candida albicans*, which method comprises:
 - (a) contacting a compound to be tested with one or more Candida albicans cells having a mutation in a nucleic acid molecule according to any of claims 1 to 8 which

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mutation results in overexpression or underexpression of said polypeptides in addition to contacting one or more wild type Candida albicans cells with said compound, monitoring the growth and/or activity of

- (b) monitoring the growth and/or activity of said mutated cell compared to said wild type; wherein differential growth or activity of said one or more mutated Candida cells is indicative of selective action of said compound on a polypeptide or another polypeptide in the same or a parallel pathway.
- 25. A compound identifiable according to the 15 method of claim 24.
 - 26. A compound according to claim 25 for use as a medicament.
- 27. Use of a compound according to claim 25 in the preparation of a medicament for treating *Candida* albicans associated diseases.
- 28. A pharmaceutical composition comprising a compound according to claim 25 together with a pharmaceutically acceptable carrier, diluent or excipient therefor.
 - 29 A method of identifying DNA sequences from a cell or organism which DNA encodes polypeptides which are critical for growth or survival of said cell or organism, which method comprises:
 - (a) preparing a cDNA or genomic library from said cell or organism in a suitable expression vector which vector is such that it can either integrate into the genome in said cell or that it permits transcription

- 41 -

of antisense RNA from the nucleotide sequences in said cDNA or genomic library.

- (b) selecting transformants exhibiting impaired growth and determining the nucleotide sequence of the cDNA or genomic sequence from the library included in the vector from said transformant.
- 30. A method according to claim 29 wherein said cell or organism is a yeast or filamentous fungus.
 - 31. A method according to claim 29 or 30 wherein said cell or organism is any of Saccharomyces cerevisiae, Schizosaccharomyces pombe or Candida albicans.
 - 32. An antibody capable of binding to a polypeptide according to claim 10 or 11.
- 20 33. An oligonucleotide comprising a fragment of from 10 to 120 contiguous nucleotides of a nucleic acid molecule according to any of claims 1 to 8.
- 34. An oligonucleotide according to claim 33 comprising a fragment of from 10 to 50 contiguous nucleotides.

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F1G.1.

AACAGCTGGT 1	CTTCTGCTAA	TACATTCAAC	CCTTTCCATA	TCTATACTCC 50
AACAATATGA 51	TAACTGATGA	ACAATTGAAT	ACCATTGCAT	TGACATTTGG 100
TTTTGCTTCA 101	ATAATATTAA	TCATAATATA	TCATGCCATA	TCTACTAATG 150
TACATAAATT 151	AGAAGATGAA	ACCCCATCAT	CTTCATTTAC	CAGAACAAAT 200
ACTACTGAAA 201	CTACTGTTGC	AAGTAAGAAA	AAGAAGTAAT	AACTGATGGA 250
TTTTTCTTCC 251	TACCACCAAT	TGAATAATGC	TAGACTTGTT	GGTGTGCTAC 300
AAATATTTCA 301	AAAGAAAATA	CGAATACTTT	ATAAAATGGT	AAGAACGGAA 350
GATGGTTTCT 351	CATTTATACA	CTAAATACAA	ATCACATACA	CATACACAAA 400
CACAAATACA	TACATACACC	TATATCCCTT	TATTTGAT 438	

	FIG. 2	<i>2</i> .		
ATGTTAAAAA 1	CACTAACTCA	AACTTTACGC	TTAACTGGGA	AAGCTTTCCC 50
-	CCCCCCTTCA	телаллеста	CGCTGCCTTC	ĞÄCCGTTCTA

AAAGGTCCGT CCGGCCTTGA TCAGAACCTA CGCTGCCTTC ĞACCGTTCTA
100

AACCTCATGT CAACATTGGT ACTATTGGTC ATGTTGATCA TGGTAAAACT 150

ACATTGACTG CTGCTATCAC CAAAGTTTTA GCCGAACAAG GTGGTGCCAA
151

CTTCTTGGAT TATGGTTCTA TTGATAGAGC TCCAGAAGAA AGAGCTAGAG 250

GTATCACTAT TTCCACTGCC CACGTTGAAT ACGAAACCAA GAACAGACAC 251

TATGCCCACG TTGATTGTCC AGGACACGCT GATTATATCA AAAATATGAT 350

TACTGGTGCC GCTCAAATGG ATGGTGCTAT CATTGTTGTT GCTGCCACTG 400

ATGGTCAAAT GCCTCAAACC AGAGAACATT TGTTATTGGC CAGACAAGTT 450

GGTGTTCAAG ACTTGGTTGT GTTTGTCAAC AAAGTCGATA CTATTGATGA
500

CCCTGAAATG TTGGAATTAG TCGAAATGGA AATGAGAGAA TTGTTATCCA
501

CCTACGGTTT TGATGGTGAC AACACTCCAG TTATTATGGG ATCTGCTTTA
551

ATGGCTTTGG AAGACAAGAA ACCAGAAATT GGTAAGGAAG CTATCTTGAA 650

ATTGTTAGAT GCTGTCGATG AACACATTCC AACTCCATCA AGAGACTTGG
651

AACAACCATT TTTGTTACCA GTTGAAGACG TGTTCTCCAT CTCCGGTAGA
701

GGAACTGTTG TCACTGGTAG AGTTGAAAGA GGTGTTTTGA AGAAGGGTGA
800

AGAAATCGAA ATTGTTGGTC GTTTTGACAA ACCTTACAAG ACTACTGTTA 850

CCGGTATTGA AATGTTCAAA AAAGAATTAG ACTCTGCTAT GGCTGGTGAC 900

FIG. 2 (CONTINUED).

AACTGTGGTG 901	TTTTGTTAAG	AGGTGTTAAA	AGAGATGAAA	TCAAGAGAGG 950
TATGGTTTTG 951	GCCAAACCAG	GTACTGCTAC	TTCTCACAAG	AAGTTCTTGG 1000
CTTCCTTGTA	TATTTTGACT	TCCGAAGAAG	GTGGTCGTTC	CACTCCATTT 1050
GGTGAAGGTT 1051	ACAAGCCTCA	ATGCTTCTTC	AGAACTAACG	ATGTCACTAC 1100
CACATTTTCA 1101	TTCCCAGAAG	GAGAAGGTGT	TGATCATTCT	CAAATGATCA 1150
TGCCAGGTGA 1151	CAACATTGAA	ATGGTTGGTG	AATTGATCAA	ATCTTGTCCA
TTAGAAGTCA 1201	ACCAACGTTT	CAACTTGAGA	GAAGGTGGTA	AAACTGTTGC 1250
TACTGGTTTG 1251	ATTACCAGAA	TCATCGAATA	AACAGAATGT	GCACTGTGAA
TAATAAAAAG 1301	AAAAGAGGTA	TATATAGGTG	ACTTTGTATT	TTGTATTGAM 1350
CAATAAAATT	CTGTAAATAG	TAAGGGCCTC		

F1G. 3

GAATTCGCCCTTAAGCACTCGTTTCAACTATACATTCAGTAACAACACCCTTAATTTACCAAACTACA TTAATGGAAGTA
TTAATGGAAGTA ACACAACGGACGCAGAGTCAGACACAACCAACAGACAGTCACCGACAACTCAGACGCAAACCCAAAG
CAAAGAGGACCA CAAAGAGGACCA
CAAAGAGGACCA GAATAGGATTTGTCAATTGATTTGCTCCACGGGTCAGTTTGGCAATTATGATTTGAATATCAACGATA
AAACTATCGTAC
AAACTATEGTAC AAGGTAAAATGACGTGGTATTTTGGAAGAGACCCCAACTCAGATTTGCAAGTGGCGTCGTCGTCGAGA
ATTTCAAACAAG
CATTTTCAAATCTGGCTCAACTTCAATGATAAATCACTATGGATAAAGGACACTTCAACTAACGGGAC
ACACCTTAACAA
CAGTCGATTGGTGAAAGGATCAAACTACCTTCTTAATCAGGGTGATGAAATAGCAGTAGGGGTTGGTA
GAGACGAGGACG
TTGTGAGGTTTGTCGTTGTCTTTGGTGACAAATACAACCCGGCAAAGCTACCTGATTCGACCAACACA
ATTAAAGATGAA
GGAATATACAAAGACTTTATTGTGAAAAATGAAACGATAGGCCAAGGAGCATTTGCCACTGTGAAAAA
GGCGATTGAACG
ATCTACGGGCGAGTCGTACGCGGTGAAGATTATAAATCGAAGAAAAGCATTAAATACCGGTGGTGGAA
GTGCCATGGCAG
GAGTGGACCGTGAATTGTCCATATTAGAGCGGCTCAACCACCCAAATATAGTTGCTCTAAAAGCTTTT
TATGAAGATATG
GACAATTACTATATTGTGATGGAATTGGTGCCGGGCGGTGATTTGATGGACTTTGTGGCTGCAAACGG
TGCAATAGGAGA
AGACGCAACACAAGTGATCACGAAACAGATTCTAGAAGGAATTGCCTATGTTCATAATTTAGGAATCT
CCCATCGTGATT
TGAAGCCAGATAATATTTTGATTATGCAAGATGACCCAATACTTGTTAAAATCACCGACTTTGGATTG
GCAAAATTCAGT GACAATCTGACGTTTATGAAAACTTTTTGTGGTACATTGGCGTATGTTGCTCCCGAAGTTATCACCGG
TAAGTATGGATC ATCGCAGATGGAACTGCAACAAAAGGACAACTACTCTTCCTTGGTTGACATTTGGTCTTTGGGATGTT
TGGTTTATGTAC
TTTTAACTTCTCATTTACCATTCAACGGGAAAAACCAGCAACAAATGTTTGCCAAGATCAAAAGGGGG
GAATTTCATGAG
GCTCCATTAAATTCATACGACATTTCTGAAGACGGAAGAGATTTCTTGCAGTGCTGCCTACAGGTTAA
TCCTAAACTAAG
GATGACGGCTGCTGAAGCTTTGAAACATAAATGGTTGCAAGACTTGTATGAAGAGGATTCTGTCAAA
CATTGAGTTTAT
CGCAATCACAGTCGCAACAATCTCGAAAGATAGATAATGGTATCCATATCGAATCATTGAGCAAAATT
CATGAAGACGTT
ATGCTTCGTCCATTGGATAGCGAAAGAAATAGGAAATCAAGTAAACAGCAAGATTTCAAGGTACCCAA
GCGTGTGATTCC
GTTATCTCAACATCCTGCAACACCGTTACCAATGTCACAACCGAAAAAGAGGCCGTATCAAATAGACC
CTAGAACAACA
AAAAAGTCGATTTGGAAGAACCTCTGACAAGCAAGAAAGTCAAGCTAAGTGATTCCGTTGTTGCGGA
GACTACTTGAAG
TTGGGGCCACTTGCAAATTCGTTATTCCAAGAAACAATAAATA
AAGAAATGACAC
TTGTGATTGCGAGATAGACGACGACAGACTATCCAAACTTCATTGTGTCATTACCAAAGAAAACGAC
CTATATGGTTAT
TGGATAAGAGTACTACTCGTGCTTGGTCAACAATACTAGTGTTGGAAAAGGCAACAAAGTTTTGCT
AGAGGAGGGAG ATATTACATCTCTTCTTTGACCCATTGTCACTGCAACATATAGGTTTCAAAGTAGTCCTTGTTGATC
ATATTACATUTUTTCITIGACCCATIGICACIGCAACATATAGGITICAAAGAAGATATAGGI
ACTGTCTGGTGA ACATAAGAGTCAAGTGGAGGTTTTGAAACAAACCTCAGAAGAAATGAATATTATTCCACTTATTTCT
ACATAAGAGTCAAGTGGAGGTTTTGAAACAAACCTCAGAAGAAATGAATATTATTGGAGTTTTAAGTAGTA
TAAGTTCATAGATTTAGCATATATACAAGCATTTCCTATAGAAACAAAGGTTCATTAATTTAGTTAT
TACCTCCATGCA
*1988 * A.M. 1 - A.M. 1

ATTACATTTACTTCTTCCCAAGGGCGAATTCTGCAGATATC

F16.4.

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/	/	CT.	S	フ.

770.0.
ATGGGTACTA GTACAAGTGA AGCATTGAAG AACATCAAAA ACAAACAGCG 50
AAGACAGAAA GTTTTTGCAG AAATAAAACA TGAAAAGAAT AAA @A ACGTC 100
ATAAGCAAAG AGCCGAAAGA GCTAAGGAAG AAAGAGAAAA CCCCGAATTA 150
AGAGAGGAAA GAATAGCAGC TAATATCCCA GATACTATAG ATAGCAAACG 200
TATTTATGAT GAGACTATAG CTGCTGAAGT TGAAGGAGAT GACGAGTTTC 250
AGTCATATTT CACTAACTTG TTGGAAGAAC CAAAGATTTT GTTGACAACA 300
AGTGCCAATG CTAAAAAACC GGCCTATGAA TTTGCAGACA TGATCATGGA 350
CTTTTTACCG AATGTGACAT TTATCAAAAG GAAGAAGGAA TATACAATGC 400
AAGATATGGC CAAATATTGC TCGAATAGAG ACTTCACTGC ATTGCTTGTC 450
ATCAACGAAG ACAAGAAGAA GGTCAATGGT ATAACGCTCA TCAATTTACC 500
TGAAGGGCCA ACATTTTATT TTTCGATTAC ATCAATAGTT GATGGGAAAA 550
GAATTAAGGG ACACGGGAAA GCTGGTGATT ATTTACCTGA GATTGTATTG 600
AATAATTTCA ATTCAAGATT GGGTAAAACT GTGGGAAGAC TATTTCAAAG 650
TATTTTCCCT CATAAACCTG AACTTCAAGG AAGACAAGTG ATTACTTTGC 700
ACAATCAACG TGATTATATT TTTTTCAGAA GACATAGATA TATTTTCAGA 750
AATGAGGAAA AGGTTGGATT GCAGGAATTG GGTCCGCAGT TTACATTAAA 800
GCTAAGAAGA ATGCAAAAGG GAGTACGTGG TGATGTTGTT TGGGAACACA 850
GACCAGATAT GGAAAGAGAT AAGAAGAAGT TTTATTTATA AGCGGGTGTA 900

FIG. 5 (CONTINUED).

TAAAGGTAGT AGTAGTGCGT TTATAAGTAT GTGTGTGTGT TTATGCATAG 901

ATGTGTAAAG AGTAATACAG CTAATTCG

F1G. 6.

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AACTAATTTG	TTTAAACATC	AATACCAAGA	AGTTTTTACA	ATTCAATCCC 50
ACATACACCA	TTAATTATGA	ATTCTGAAAA	GATTATTGAA	GTTATCATTG 100
CTATTTTCTT	ACCACCAGTA	GCTGTGTTTA	TGAAATGTGG	TGCCACTACC 150
CCATTATGGA	TTAACTTGGT	ATTATGTATC	TTTATTTGGT	TCCCTGCTAT 200
CTTACATGCC	TTATACGTTG	TGTTGAAAGA	TTAAACAAAC	ACCAGAGATT 250
TACTGCTTGA	TGAATTGATT	ACTCCAAAGA	GTTGTGACTA	GTTCCCAGTG 300
- · -	GCCTTCCAAC	TTTCTTTTAC	ATTTTTCCAT	TACTACCACT 350
	CTATTTTGCA	GAGTTTTCAA	AATTTATCCA	AAACATGTTA 400
	CATATTATT	A TAATTATTCT	TTTTTGTATI	TTTTTCCCTT 450
	ATTTATTA	A TCGTTTCGT	GTTTGGTAT	TTATTTTTTT 500
	A ATTCGAATA	r atatctatac	2 ATGAATTTA	TATCCATTGT 550
	T AAAACATTT	T GTTAGTTTT	T TGTTACTAG	TANNAAANNAT 000
AATAAAAGT 601	T TANTTCAAC			

9/53 FIG. 7.

	,,,	• • •		
ATGACATTAG 1	GGTTCGATAA	ATTCATAAGC	AAGGTCAGCA	CTCATAGACG 50
TCAATCTGAA 51	CCATCAATCT	TGGAAATCGC	AGCCACCAAT	TCTCAAAATA 100
AATCGAGAAG 101	GCTAAGTATG	GATAATGGTC	ATTGTTATGT	TCGTGAATCA 150
ACTAATAATC 151	ATCATCATTT	AAATACCGTC	GTTGATAATT	TACGACAGCG 200
TGCGGGATCG 201	TTTTCATTTA	TTTCACATCA	CCATAATCAC	CATCAGAATA 250
GTCACGATAA 251	TTATACTGTC	GATCCCCTTA	CATCAAACGG	AGCACGAATT 300
TCCCGATCAC 301	GTTCACGTTC	CAAATCAGTT	GGGCACGGAG	AAGCAATATC 350
ACCAGCGTAT 351	TTTTCCAAGA	ATAAAACCAA	AGATTTAGTG	AAACAGGAAA 400
CAGCACATAT 401	CATTCTGAAG	AAATTACTCA	ACATGTTACA	AGATTTGGAT 450
TTACAAAACC 451	CTATTGCATT	GAAAACAATA	TCACAAGGTT	CAGAATCAAA 500
GTTTTGTAAA 501	ATCTACGTGT	CTAACACTAA	TAATTGTATT	TACTTACCAG 550
CAGCAAGTTC 551	AACAAGTTTC	ACTTATGAAG	ATGATGAAAA	TGGCGGCGTT 600
ATAATTGCTG 601	AAGATAGAAA	TGATGAAATG	CCAACAGCAG	TTAATAACAA 650
TACTTTGTCA 651	ATGGATAGTA	TAAATCATIC	AGAGACTGAT	TTCCTGGATT 700
CTCCACCACC 701	TCCAGATTTA	TTTTCTAAAA	TGAAATCATI	CCATTCACCA 750
AATTACTTGA 751	CTTCAAAAAT	CGATTCTGAA	TGTCCAATTC	CACATACATT
TGCTGTGATT 801	GTTGAATTAA	. CCAAGGACTO	TTTGATTATT	AAAGATCTTC 850
ATTTCCAATT	TCAGTCATTA	ACTACCATEI	TATEGECAAC	TGGGGATGCA

F16	TICONTIN	WED1).		r gujaș
TATAATCGGA 901	CTCATGCCAA	GGAGAAATTT	ACCATTGGGA	ATATGGAATG 950
GCGTACATCT 951	TTAAGCGACG	CCGACTATTA	TATCAATAGT	TCTAATTCCA 1000
ACGATGTTAA 1001	GCTGAAAAAC	TTGGGTCCTG	AAGATCTTAT	TAATCGAACT 1050
AGAGAATACA 1051	AATTAATCGA	TATTGAAGAA	CCAAACAATT	CATCAAACAG 1100
TTTACTGGAT	GATGACATGG	ATATTAATAA	TATTACGTCG	CCATTATCAA 1150
CGTCACCAAC 1151	ATCAAGTTCA	ACTTCAACAA	ATTCAACCTC	CAACTCATTG 1200
GGTTCAGATT 1201	CATATAAAGC	TGGTCTTTAT	GTATTTTTAT	TACCAATCTT 1250
ATTGCCAGAA 1251	CATATTCCTG	CTTCCATTGT	TTCTATTAAT	GGTTCATTGG 1300
CTCATACATT	ACTGGTTGAA	TGCAATAAAT	ATACTGATAA	GTTGAATCGG 1350
AAATCAAAAG 1351	TATCAGCATC	GTACAATTTA	CCTATGGTCC	GTACTCCACC 1400
AAACATTGGT 1401	AATTCCATTG	CTGATAAGCC	AATTTATGTT	AATAGGATTT 1450
GGAATGATGC 1451	CGTACATTAT	ATTATAACTT	TCCCCCGCAA	ATATGTTACT 1500
TTGGGTTGTG 1501	AACACATGAT	AAATGTGAAA	TTACTGCCCA	TGGTGAAAGA 1550
TGTGGTTATC 1551	AAGCGTATTA	AATTTAAIGT	ATTGGAGAGA	ATAACTTATG 1600
TTTCCAAAAA 1601	TTTATCACGA	GAATATGAIT	ATGATAGTGA	AGACCCCTAT 1650
TGTATTCATC 1651	CAGTTICTAA	AGAAAATAAA	GTACGTGAAC	GTGTTGTGTC 1700
GTTATATGAA 1701	TTGAAAACGA	AGGCAAAACA	ATCTTCTGGT	GGACATCTTG 1750
AAGCTTATAA 1751	ACAAGAAGTT	ATGAAATGTC	CGGAAAATAA	CCTTTTATTT 1800

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TCTTGTTATG 1801	AGGTTGAAAA	TGATAATAAT	AACGGCAACG	GCAACGGCAA 1850
CGGCAACGGA 1851	AACAAGAACG	TTAAACAAAA	GAATAAAGAT	CAACCAATGA 1900
TTGCTACACC 1901	TTTAGATATC	AATGTTTCTT	TACCATTTTT	AACTACTATG 1950
TCTGATAGTT 1951	TAATTATGAC	ATCAGCCATA	GAAGAAGAAG	GTTCAGATCT 2000
GCCTCATACA 2001	TCAAGAAGAG	GGTCGGCAGT	GAGTATGACT	GATAATAATA 2050
CTACCCCAAG 2051	TAACAATAAC	CCTTTATCTC	CATTTTTGGG	AGCAGTGGAA 2100
ACTAATGGTG 2101	CTAGTATAAA	TGAAATTGGT	GATCATACAT	TATTCCCTGA 2150
TTCTAATTTT 2151	CGACATATTG	AAATTAAACA	TCGATTACAA	GTTACATTTA 2200
GGATTTCTAA 2201	ACCGGATCTG	GATAATAAAA	TGCATCATTA	TGAAGTGGTT 2250
ATTGATACCC 2251	CCATCGTTTT	ACTTAGTTCA	AAATGTCAAG	AAGATTCTCC 2300
TCCTCCTTAT 2301	AGTTCTGTA 2319			

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12/53

F16.8.

AACGTTCGTG CAAAAGGCTA TACTGGTGAT ATCCACGCAG ATGAAGAGCA 50

AGTTTAATCA ACTCTTTGTC AATTAATGCT GTACTTGTTT TCATTTTATT 100

TGCTGGCATT TAAAGAATAC CCATAGTTCA GAAAATAAAA TTGAAAAAATT 151

TAAAAAAAAA CGCAATATCA TTCATTTTTT TTGTTTTTTT GACAATAATA 200

TTAATATGTA GTTACCAATG TTTTTAGATT TTATATGTTT TGAAAAAATA 250

GTTTG

F1G. 9.

AACCTTACAA TCATTATACC AACTATCAAA ATCATAAGAC TCTTNAACTT
1 50
CTGTTTTTGA TAGTTGGTAT AATGATTTAT GTATTATCTT AATTCATTAT
51 100
TATTAGTTTC GGTCACAAA
101 119

PCT/EP99/09833

13/53 FIG. 10. CYCT BamH I (6901) Ava I (6842) Pst I (6635) **HIFN**B Hind III (1) ORI Ava I (6067) Apa LI (704) BamH I (6035) Ava I (6029) Pst I (6021) EcoR I (5704) AMP Hind III (5403) pGAL1PNiST-1 Pgal1 7177 bp Apa LI (1950) Hind III (5363) Apa LI (4999) EcoR I (4957) Apa LI (2447)

CaURA3

EcoR I (3827)

14/53 FIG. 11.

1 AGCTTGAGTA TTCTATAGTG TCACCTAAAT AGCTTGGCGT AATCATGGTC 51 ATAGCTGTTT CCTGTGTGAA ATTGTTATCC GCTCACAATT CCACACAACA 101 TACGAGCCGG AAGCATAAAG TGTAAAGCCT GGGGTGCCTA ATGAGTGAGC 151 TAACTCACAT TAATTGCGTT GCGCTCACTG CCCGCTTTCC AGTCGGGAAA 201 CCTGTCGTGC CAGCTGCATT AATGAATCGG CCAACGCGCG GGGAGAGGCG 251 GTTTGCGTAT TGGGCGCTCT TCCGCTTCCT CGCTCACTGA CTCGCTGCGC 301 TCGGTCGTTC GGCTGCGGCG AGCGGTATCA GCTCACTCAA AGGCGGTAAT 351 ACGGTTATCC ACAGAATCAG GGGATAACGC AGGAAAGAAC ATGTGAGCAA 401 AAGGCCAGCA AAAGGCCAGG AACCGTAAAA AGGCCGCGTT GCTGGCGTTT 451 TTCCATAGGC TCCGCCCCC TGACGAGCAT CACAAAAATC GACGCTCAAG 501 TCAGAGGTGG CGAAACCCGA CAGGACTATA AAGATACCAG GCGTTTCCCC 551 CTGGAAGCTC CCTCGTGCGC TCTCCTGTTC CGACCCTGCC GCTTACCGGA 601 TACCTGTCCG CCTTTCTCCC TTCGGGAAGC GTGGCGCTTT CTCATAGCTC 651 ACGCTGTAGG TATCTCAGTT CGGTGTAGGT CGTTCGCTCC AAGCTGGGCT 701 GTGTGCACGA ACCCCCCGTT CAGCCCGACC GCTGCGCCTT ATCCGGTAAC 751 TATCGTCTTG AGTCCAACCC GGTAAGACAC GACTTATCGC CACTGGCAGC 801 AGCCACTGGT AACAGGATTA GCAGAGCGAG GTATGTAGGC GGTGCTACAG 851 AGTTCTTGAA GTGGTGGCCT AACTACGGCT ACACTAGAAG GACAGTATTT 901 GGTATCTGCG GTCTGCTGAA GCCAGTTACC TTCGGAAAAA GAGTTGGTAG 951 CTCTTGATCC GGCAAACAAA CCACCGCTGG TAGCGGTGGT TTTTTTGTTT 1001 GCAAGCAGCA GATTACGCGC AGAAAAAAAG GATCTCAAGA AGATCCTTTG 1051 ATCTTTTCTA CGGGGTCTGA CGCTCAGTGG AACGAAAACT CACGTTAAGG 1101 GATTTTGGTC ATGAGATTAT CANAAAGGAT CTTCACCTAG ATCCTTTTAA 1151 ATTAAAAATG AAGTTTTAAA TCAATCTAAA GTATATATGA GTAAACTTGG 1201 TCTGACAGTT ACCAATGCTT AATCAGTGAG GCACCTATCT CAGCGATCTG 1251 TCTATTTCGT TCATCCATAG TTGCCTGACT CCCCGTCGTG TAGATAACTA 1301 CGATACGGGA GGGCTTACCA TCTGGCCCCA GTGCTGCAAT GATACCGCGA

FIG. 11 (CONTINUED 1).

1351 GACCCACGCT CACCGGCTCC AGATTTATCA GCAATAAACC AGCCAGCCGG 1401 AAGGGCCGAG CGCAGAAGTG GTCCTGCAAC TTTATCCGCC TCCATCCAGT 1451 CTATTAATTG TTGCCGGGAA GCTAGAGTAA GTAGTTCGCC AGTTAATAGT 1501 TTGCGCAACG TTGTTGCCAT TGCTACAGGC ATCGTGGTGT CACGCTCGTC 1551 GTTTGGTATG GCTTCATTCA GCTCCGGTTC CCAACGATCA AGGCGAGTTA 1601 CATGATCCCC CATGTTGTGC AAAAAAGCGG TTAGCTCCTT CGGTCCTCCG 1651 ATCGTTGTCA GAAGTAAGTT GGCCGCAGTG TTATCACTCA TGGTTATGGC 1701 AGCACTGCAT AATTCTCTTA CTGTCATGCC ATCCGTAAGA TGCTTTTCTG 1751 TGACTGGTGA GTACTCAACC AAGTCATTCT GAGAATAGTG TATGCGGCGA 1801 CCGAGTTGCT CTTGCCCGGC GTCAATACGG GATAATACCG CGCCACATAG 1851 CAGAACTTTA AAAGTGCTCA TCATTGGAAA ACGTTCTTCG GGGCGAAAAC 1901 TCTCAAGGAT CTTACCGCTG TTGAGATCCA GTTCGATGTA ACCCACTCGT 1951 GCACCCAACT GATCTTCAGC ATCTTTACT TTCACCAGCG TTTCTGGGTG 2001 AGCAAAAACA GGAAGGCAAA ATGCCGCAAA AAAGGGAATA AGGGCGACAC 2051 GGAAATGTTG AATACTCATA CTCTTCCTTT TTCAATATTA TTGAAGCATT 2101 TATCAGGGTT ATTGTCTCAT GAGCGGATAC ATATTTGAAT GTATTTAGAA 2151 ARATARACAR ATAGGGGTTC CGCGCACATT TCCCCGAAAA GTGCCACCTG 2201 ACGTCTAAGA AACCATTATT ATCATGACAT TAACCTATAA AAATAGGCGT 2251 ATCACGAGGC CCTTTCGTCT CGCGCGTTTC GGTGATGACG GTGAAAACCT 2301 CTGACACATG CAGCTCCCGG AGACGGTCAC AGCTTGTCTG TAAGCGGATG 2351 CCGGGAGCAG ACAAGCCCGT CAGGGCGCGT CAGCGGGTGT TGGCGGGTGT 2401 CGGGGCTGGC TTAACTATGC GGCATCAGAG CAGATTGTAC TGAGAGTGCA 2451 CCATATGCGG TGTGAAATAC CGCACAGATG CGTAAGGAGA AAATACCGCA 2501 TCAGGCGAAA TTGTAAACGT TAATATTTTG TTAAAATTCG CGTTAAATAT 2551 TTGTTAAATC AGCTCATTTT TTAACCAATA GGCCGAAATC GGCAAAATCC 2601 CTTATAAATC AAAAGAATAG ACCGAGATAG GGTTGAGTGT TGTTCCAGTT 2651 TGGAACAAGA GTCCACTATT AAAGAACGTG GACTCCAACG TCAAAGGGCG

FIG. HICONTINUED 2).

2701 AAAAACCGTC TATCAGGGCG ATGGCCCACT ACGTGAACCA TCACCCAAAT 2751 CAAGTTTTTT GCGGTCGAGG TGCCGTAAAG CTCTAAATCG GAACCCTAAA 2801 GGGAGCCCCC GATTTAGAGC TTGACGGGGA AAGCCGGCGA ACGTGGCGAG 2851 AAAGGAAGGG AAGAAAGCGA AAGGAGCGGG CGCTAGGGCG CTGGCAAGTG 2901 TAGCGGTCAC GCTGCGCGTA ACCACCACAC CCGCCGCGCT TAATGCGCCG 2951 CTACAGGGCG CGTCCATTCG CCATTCAGGC TGCGCAACTG TTGGGAAGGG 3001 CGATCGGTGC GGGCCTCTTC GCTATTACGC CAGCTGGCGA AAGGGGGATG 3051 TGCTGCAAGG CGATTAAGTT GGGTAACGCC AGGGTTTTCC CAGTCACGAC 3101 GTTGTAAAAC GACGGCCAGT GAATTGTAAT ACGACTCACT ATAGGGCGAA 3151 TTGGTTTTCC AATGATGAGC ACTTTTAAAG TTCTGCTATG TGGCGCGGTA 3201 TTATCCCGTG TTGACGCCGG GCAAGAGCAA CTCGGTCGCC GCATACACTA 3251 TTCTCAGAAT GACTTGGTTG AGTACTAATA GGAATTGATT TGGATGGTAT 3301 AAACGGAAAC AAAAAAAAGA GCTGGTACTA CTTTCTTTAA AATTATTTTA 3351 TTATTTGATT TTATTTAATA GTATATATTA TATTTTGAAC GTAGATTATT 3401 TTGTTGAAAG TTGCTGTAGT GCCATTGATT CGTAACACTA ATTCTGTATT 3451 AGTCATTCCT CTTGTTTGAT AGTATCCAAA AAAACGGCTA TTTTTTTGCA 3501 ATCTTATTTC CTGCATATTA TACAGATAAC ATAATGAAAG AAAAAATCTT 3551 TTTTTTGTT CTTCAATGAT GATTTCAACC ATTCTTTAA ACATTGATCA 3601 ATTCCTGAGC AACAACCCCA TACACACTGG TTTATATACC GCCCCTTTTA 3651 CAGTTGAAGA AAGAAATAGA AATAGAAATA GCAAACAAAA GATATGACAG 3701 TCAACACTAA GACCTATAGT GAGAGAGCAG AAACTCATGC CTCACCAGTA 3751 GCACAGCGAT TATTTCGATT AATGGAACTG AAGAAAACCA ATTTATGTGC 3801 ATCAATTGAC GTTGATACCA CTAAGGAATT CCTTGAATTA ATTGATAAAT 3851 TAGGTCCTTA TGTATGCTTA ATCAAGACTC ATATTGATAT AATCAATGAT 3901 TTTTCCTATG AATCCACTAT TGAACCATTA TTAGAACTTT CACGTAAACA 3951 TCAATTTATG ATTTTTGAAG ATAGAAAATT TECTGATATT GGTAATACCG 4001 TAAAGAAACA ATATATTGGT GGAGTTTATA AAATTAGTAG TTGGGCAGAT

FIG. HI (CONTINUED 3).

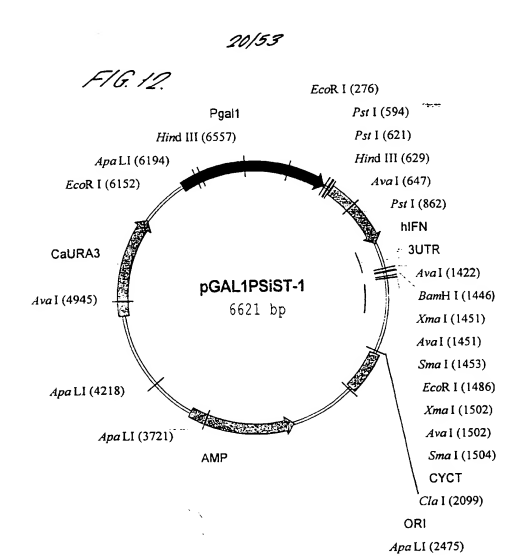
4051 ATTACCAATG CTCATGGTGT CACTGGGAAT GGAGTGGTTG AAGGATTAAA 4101 ACAGGGAGCT AAAGAAACCA CCACCAACCA AGAGCCAAGA GGGTTATTGA 4151 TGTTAGCTGA ATTATCATCA GTGGGATCAT TAGCATATGG AGAATATTCT 4201 CAAAAAACTG TTGAAATTGC TAAATCCGAT AAGGAATTTG TTATTGGATT 4251 TATTGCCCAA CGTGATATGG GTGGCCAAGA AGAAGGATTT GATTGGCTTA 4301 TTATGACACC TGGAGTTGGA TTAGATGATA AAGGTGATGG ATTAGGACAA 4351 CAATATAGAA CTGTTGATGA AGTTGTTAGC ACTGGAACTG ATATTATCAT 4401 TGTTGGTAGA GGATTGTTTG GTAAAGGAAG AGATCCAGAT ATTGAAGGTA 4451 AAAGGTATAG AAATGCTGGT TGGAATGCTT ATTTGAAAAA GACTGGCCAA 4501 TTATAAATGT GAAGGGGGAG ATTTTCACTT TATTAGATTT GTATATATGT 4551 AGAATAAATA AATAAATAAG TTAAATAAAT AATTAAATAA GGGTGGTAAT 4601 TATTACTATT TACAATCAAA GGTGGTCCTT CTAGCTGTAA TCCGGGCAGC 4651 GCAACGGAAC ATTCATCAGT GTAAAAATGG AATCAATAAA GCCCTGCGCA 4701 GCGCGCAGGG TCAGCCTGAA TACGCGTTTA ATGACCAGCA CAGTCGTGAT 4751 GGCAAGGTCA GAATAGCCCA AGTCGGCCGA GGGGCCTGTA CAGTGAGGGA 4801 AGATCTGATA TTGACGAAGA GGAACCAATG TAACGTTACA CTGAAGAAAA 4851 CACATAATAA ACGGGAAGAA ACGGTGTAAA AGTGTGAAAA TAATTTTTGA 4901 ATATCATTTC CCTTGGTTTA ATTCCAAACG AAACGTGTAT TTTTTTAGAG 4951 AATGGGAATT CTTATTGGAT GTCTAGATTG TTTGTTTACT CCAGACTGTG 5001 CACAAAAACG TTTGGATGGA TGATCAGAAG ATATTTTTAG GCTTAGCTCT 5051 AAATATAAGA AATGATGCTT GAAAATCCAG ACAGAAATTG AGTTTCAAAA 5101 ATTGGTAATG TGAGGTATTA GTCAACTAAC CAAATAACAA TGCAAACCGG 5151 TTGATACATT TCATTTTGAA AATAATGAAA CTGGAATTGG ATGACCAGCA 5201 CACAAACACA TAAAGTAATT ATGGGAATTA GAAGCGAACA TAGAGGAATA 5251 CTTTGCCACG AACAGAATAC AAGTGGGAAC ACTTTTTTCT CCATTGTTTT 5301 AGTTCTGTTT TTTTGTCAAA CTGGTTTTGT GCTATGTGTA AAAAAATATT 5351 GCCAAGAAA AAAGCTTGTT TTGTGGCCAG TGTCCGAAAA AAATTTTGGG

FIG. H (CONTINUED 4).

5401	GAAGCTTCGG	TATTTAT	TTTTTTTTTC	CATCGGGGAA	AGTGGGGGG
5451	TTAAAAAAA	TAAGCAGTTC	ATAAAACCTT	ССААААААТА	TATGGACAGA
5501	GATGATTGTA	TTTTCCCGAC	ACCAAAATCA	TAATTAACTA	TGAGAAAATT
5551	GAATGTAACG	TTACAATTTA	TTTTTATTTG	AAGCTGAAAA	GCGATTTATG
5601	ATTTTTCCGA	AATGAAAATT	TTTTTTAGGT	TTATTTTTTT	TGTCGGGCAA
5651	AGAAAAACTG	AACAAGGATT	ATTAAAATTT	TTGGTGTTTG	TTTGTGTCTG
5701	GAGAATTCAT	TCCTCTCTCA	TCTTCACACA	ATGTTTAGAC	ATCTGACACG
5751	ATTCAAAATA	GTTCGGTTTC	CGGGGTTGGT	GTTTAGTTTT	CGTTTTTCGT
5801	TTTTTTTGGA	AAGAATGTTT	TAGCTCATTG	GTTTTCTTTC	TTCATTCAAT
5851	AGTTTTGAAA	GAATTTGCCC	ACTTGTTATT	ACAATCATAT	AAAATTAAAC
5901	TTTGATATAA	AATAGAGTTT	GAAAGTTTCC	CAGATCCTTT	TTGATTTCTT
5951	TGTAATTTTT	TTTTCTCCCA	CATATACACA	CATACAAACC	GATTTTTATA
6001	AGAAAGAGTT	ATACCCTGCA	GCTCGACCTC	GAGGGATCCG	GGCCCTCTAG
6051	ATGCGGCCGC	TAGGCCTCGA	GGGACTTTTG	CACCAAAAAT	AATTTATTTT
6101	CCAAAATAAA	AŢTTAAATAA	АТААААТА	CTCATAATTT	AATAAAAATT
6151	TCAAAATCTT	CTAGTGTCCT	TTCATATGCA	GTACATTAGC	CATCAGTCAC
6201	TTAAACAGCA	TCTGCTGGTT	GAAGAATGCT	TGAAGCAATT	GTCCAGTCCC
6251	AGAGGCACAG	GCTAGGAGAT	CTTCAGTTTC	GGAGGTAACC	TGTAAGTCTG
6301	TTAATGAAGT	AAAAGTTCCT	TAGGATTTCC	ACTCTGACTA	TGGTCCAGGC
6351	ACAGTGACTG	TACTCCTTGG	CCTTCAGGTA	ATGCAGAATC	CTCCCATAAT
6401	ATCTTTTCAG	GTGCAGACTG	CTCATGAGTT	TTCCCCTGGT	GAAATCTTCT
6451	TTCTCCAGTT	TTTCTTCCAG	GACTGTCTTC	AGATGGTTTA	TCTGATGATA
6501	GACATTAGCC	: AGGAGGTTCT	CAACAATAGI	CTCATTCCAG	CCAGTGCTAG
6551	ATGAATCTTG	TCTGAAAATA	GCAAAGATG1	TCTGGAGCAT	CTCATAGATG
6601					TAATCTCCTC
6651					A AGCCTCCCAT
6701					TCTTTGTAGG

FIG. 11 (CONTINUED 5).

AATCCAAGCA AGTTGTAGCT CATGGAAAGA GCTGTAGTGG AGAAGCACAA
6801 CAGGAGAGCA ATTTGGAGGA GACACTTGTT GGTCATGTTC CTCGAGGCCT
6851 TTTTGGCCAG CTGGCGCCTG CTGCGCGACG GCGAGCTGCT CACCACCCAG
6901 GATCCGTCCC CCTTTTCCTT TGTCGATATC ATGTAATTAG TTATGTCACG
6951 CTTACATTCA CGCCCTCCCC CCACATCCGC TCTAACCGAA AAGGAAGGAG
7001 TTAGACAACC TGAAGTCTAG GTCCCTATTT ATTTTTTAT AGTTATGTTA
7051 GTATTAAGAA CGTTATTTAT ATTTCAAATT TTTCTTTTTT TTCTGTACAG
7101 ACGCGTGTAC GCATGTAACA TTATACTGAA AACCTTGCTT GAGAAGGTTT
7151 TGGGACGCTC GAAGGCTTTA ATTTGCA



F1G. 13.

1	TTCCATCGGG	GAAAGTGGGG	GGGAAAAAAT	TTTAAGCAGT	TCACAAAACC
51	TTCCAAAAAA	TATATGGACA	AAGATGATTG	TATTTTCCCG	ACACCAAAAT
101	CATAATTAAT	TATGAGAAAG	TTAAATGTAA	CGTTACAATT	TATGTTTATT
151	TGAAGGTGAA	AAGCGATTTA	TGATTTTTCC	GAAATGAAAA	TTTTTTTAG
201	GTTTATTTTT	TTTGTCGGGC	AAAGAAAAAC	TGAACAAGGA	TTATTAAAAT
251	TTTTGGTGTT	TGTTTGTGTC	TGGAGAATTC	ATTCCTCTCT	CATCTTCACA
301	CAATGTTTAG	ACATCTGACA	CGATTCATGA	TAGTTCGGTT	TCCGGGGTTG
351	GTGTTTAGTT	TTCGTTTTTC	TTTTTTTTTG	GAAAGAATGT	TTTAGCTCAT
401	TGGTTTTCTT	TCTTCATTCA	ATAGTTTTGA	AAGAATTTGC	CCACTTGTTA
451	TTACAATCAT	ATAAAATTAA	ACTTTGATAT	AAAATAGAGT	TTGAAAGTTT
501	CCCAGATCCT	TTTTGATTTC	TTTGTAAATT	TTTTTTTCTC	CCACATATAC
551	ACACATACAA	ACCGATTTTT	ATAAGAAAGA	GTTATACCCT	GCAGCTCGAC
601	CTCGACTGTT	TAAACCTGCA	GGCATGCAAG	CTTGGCCAAA	AAGGCCTCGA
651	GGAACATGAC	CAACAAGTGT	CTCCTCCAAA	TTGCTCTCCT	GTTGTGCTTC
701	TCCACTACAG	CTCTTTCCAT	GAGCTACAAC	TTGCTTGGAT	TCCTACAAAG
751	AAGCAGCAAT	TTTCAGTGTC	AGAAGCTCCT	GTGGCAATTG	AATGGGAGGC
801	TTGAATACTG	CCTCAAGGAC	AGGATGAACT	TTGACATCCC	TGAGGAGATT
851	AAGCAGCTGC	AGCAGTTCCA	GAAGGAGGAC	GCCGCATTGA	CCATCTATGA
901	GATGCTCCAG	AACATCTTTG	CTATTTCAG	ACAAGATTCA	TCTAGCACTG
951	GCTGGAATGA	GACTATTGTT	GAGAACCTCC	TGGCTAATGT	CTATCATCAG
1001	ATAAACCATC	TGAAGACAGT	CCTGGAAGAA	AAACTGGAGA	AAGAAGATTT
1051	CACCAGGGGA	AAACTCATGA	GCAGTCTGCA	CCTGAAAAGA	TATTATGGGA
1101	GGATTCTGCA	TTACCTGAAG	GCCAAGGAGT	ACAGTCACTG	TGCCTGGACC
1151	ATAGTCAGAG	TGGAAATCCT	AAGGAACTTT	TACTTCATTA	ACAGACTTAC
1201	AGGTTACCTC	: CGAAACTGAA	GATCTCCTAG	CCTGTGCCTC	TGGGACTGGA
1251	CAATTGCTTC	: AAGCATTCTI	CAACCAGCAG	ATGCTGTTTA	AGTGACTGAT
1301	GGCTAATGT	CTGCATATGA	AAGGACACTA	GAAGATTTT	AAATTTTTAT

FIG. 13 (CONTINUED 1).

1401 TATTTTTGGT GCAAAAGTCC CTCGAGGCCT AGCGGCCGCC TAGAGGATCC 1451 CCGGGCGCTA GGCGGCCGCT AGGCCTTTTT GGCCGAATTC GAGCTCGGTA 1501 CCCGGGGAGA TCCGTCCCCC TTTTCCTTTG TCGATATCAT GTAATTAGTT 1551 ATGTCACGCT TACATTCACG CCCTCCCCCC ACATCCGCTC TAACCGAAAA 1601 GGAAGGAGTT AGACAACCTG AAGTCTAGGT CCCTATTTAT TTTTTTATAG 1651 TTATGTTAGT ATTAAGAACG TTATTTATAT TTCAAATTTT TCTTTTTTTT 1701 CTGTACAGAC GCGTGTACGC ATGTAACATT ATACTGAAAA CCTTGCTTGA 1751 GAAGGTTTTG GGACGCTCGA AGGCTTTAAT TTGCAAGCTA GCTTGGCGTA 1801 ATCATGGTCA TAGCTGTTTC CTGTGTGAAA TTGTTATCCG CTCACAATTC 1851 CACACAACAT ACGAGCCGGA AGCATAAAGT GTAAAGCCTG GGGTGCCTAA 1901 TGAGTGAGCT AACTCACATT AATTGCGTTG CGCTCACTGC CCGCTTTCCA 1951 GTCGGGAAAC CTGTCGTGCC AGAGATCTCT GCATTAATGA ATCGGCCAAC 2001 GCGCGGGGAG AGGCGGTTTG CGTATTGGGC GCTCTTCCGC TTCCTCGCTC 2051 ACTGACTCGC TGCGCTCGGT CGTTCGGCTG CGGCGAGCGG TATCAGATCG 2101 ATCTCACTCA AAGGCGGTAA TACGGTTATC CACAGAATCA GGGGATAACG 2151 CAGGAAAGAA CATGTGAGCA AAAGGCCAGC AAAAGGCCAG GAACCGTAAA 2201 AAGGCCGCGT TGCTGGCGTT TTTCCATAGG CTCCGCCCCC CTGACGAGCA 2251 TCACAAAAT CGACGCTCAA GTCAGAGGTG GCGAAACCCG ACAGGACTAT 2301 AAAGATACCA GGCGTTTCCC CCTGGAAGCT CCCTCGTGCG CTCTCCTGTT 2351 CCGACCCTGC CGCTTACCGG ATACCTGTCC GCCTTTCTCC CTTCGGGAAG 2401 CGTGGCGCTT TCTCATAGCT CACGCTGTAG GTATCTCAGT TCGGTGTAGG 2451 TCGTTCGCTC CANGCTCGGC TCTGTGCACG AACCCCCCGT TCAGCCCGAC 2501 CGCTGCGCCT TATCCGGTAA CTATCGTCTT GAGTCCAACC CGGTAAGACA 2551 CGACTTATCG CCACTGGCAG CAGCCACTGG TAACAGGATT AGCAGAGCGA 2601 GGTATGTAGG CGGTGCTACA GAGTTCTTGA AGTGGTGGCC TAACTACGGC 2651 TACACTAGAA GCACAGTATT TGGTATCTGC GCTCTGCTGA AGCCAGTTAC

FIG. 13 (CONTINUED 2).

	770.70 (20,77	
2701	CTTCGGAAAA AGAGTTGG	GTA GCTCTTGATC CGGCAAACAA ACCACCGCTG
2751	GTAGCGGTGG TTTTTTTG	GTT TGCAAGCAGC AGATTACGCG CAGAAAAAA
2801	GGATCTCAAG AAGATCCT	TTT GATCTTTTCT ACGGGGTCTG ACGCTCAGTG
2851	GAACGAAAAC TCACGTTA	AAG GGATTTTGGT CATGAGAFTA TCAAAAAGGA
2901	TCTTCACCTA GATCCTTT	TTA AATTAAAAAT GAAGTTTTAA ATCAATCTAA
2951	AGTATATATG AGTAAACT	TTG GTCTGACAGT TACCAATGCT TAATCAGTGA
3001	GGCACCTATC TCAGCGAT	TCT GTCTATTTCG TTCATCCATA GTTGCCTGAC
3051	TCCCCGTCGT GTAGATAA	ACT ACGATACGGG AGGGCTTACC ATCTGGCCCC
3101	AGTGCTGCAA TGATACCO	GCG AGACCCACGC TCACCGGCTC CAGATTTATC
3151	AGCAATAAAC CAGCCAGC	CCG GAAGGCCGA GCGCAGAAGT GGTCCTGCAA
3201	CTTTATCCGC CTCCATC	CAG TCTATTAATT GTTGCCGGGA AGCTAGAGTA
3251	AGTAGTTCGC CAGTTAAT	TAG TTTGCGCAAC GTTGTTGCCA TTGCTACAGG
3301	CATCGTGGTG TCACGCTC	CGT CGTTTGGTAT GGCTTCATTC AGCTCCGGTT
3351	CCCAACGATC AAGGCGAC	GTT ACATGATCCC CCATGTTGTG CAAAAAAGCG
3401	GTTAGCTCCT TGGGTCC	CTCC GATCGTTGTC AGAAGTAAGT TGGCCGCAGT
3451	GTTATCACTC ATGGTTAT	ATGG CAGCACTGCA TAATTCTCTT ACTGTCATGC
3501	CATCCGTAAG ATGCTTT	TTCT GTGACTGGTG AGTACTCAAC CAAGTCATTC
3551	TGAGAATAGT GTATGCG	GGCG ACCGAGTTGC TCTTGCCCGG CGTCAATACG
3601	GGATAATACC GCGCCAC	CATA GCAGAACTTT AAAAGTGCTC ATCATTGGAA
3651	AACGTTCTTC GGGGCGA	AAAA CTCTCAAGGA TCTTACCGCT GTTGAGATCC
3701	AGTTCGATGT AACCCAC	CTCG TGCACCCAAC TGATCTTCAG CATCTTTTAC
3751	TTTCACCAGC GTTTCTG	eggt gagcaaaaac aggaaggcaa aatgccgcaa
3801	AAAAGGGAAT AAGGGCG	GACA CGGAAATGTT GAATACTCAT ACTCTTCCTT
3851	TTTCAATATT ATTGAAG	GCAT TTATCAGGGT TATTGTCTCA TGAGCGGATA
3901	CATATTTGAA TGTATTT	TAGA AAAATAAACA AATAGGGGTT CCGCGCACAT
3951	TTCCCCGAAA AGTGCCA	ACCT GACGTCTAAG AAACCATTAT TATCATGACA
4001	TTAACCTATA AAAATAG	GGCG TATCACGAGG CCCTTTCGTC TCGCGCGTTT

FIG. 13 (CONTINUED 3). 4051 CGGTGATGAC GGTGAAAACC TCTGACACAT GCAGCTCCCG GAGACGGTCA 4101 CAGCTTGTCT GTAAGCGGAT GCCGGGAGCA GACAAGCCCG TCAGGGCGCG 4151 TCAGCGGGTG TTGGCGGGTG TCGGGGCTGG CTTAACTATG CGGCATCAGA 4201 GCAGATTGTA CTGAGAGTGC ACCATATCGA CGCTCTCCCT TATGCGACTC 4251 CTGCATTAGG AAGCAGCCCA GTAGTAGGTT GAGGCCGTTG AGCACCGCCG 4301 CCGCAAGGAA TGGTGCATGC AAGGAGATGG CGCCCAACAG TCCCCCGGCC 4351 ACGGGGCCTG CCACCATACC CACGCCGAAA CAAGCACTAA TAGGAATTGA 4401 TTTGGATGGT ATAAACGGAA ACAAAAAAA GAGCTGGTAC TACTTTCTTT 4451 AAAATTATTT TATTATTTGA TTTTATTTAA TAGTATATAT TATATTTTGA 4501 ACGTAGATTA TTTTGTTGAA AGTTGCTGTA GTGCCATTGA TTCGTAACAC 4551 TAATTCTGTA TTAGTCATTC CTCTTGTTTG ATAGTATCCA AAAAAACGGC 4601 TATTTTTTT CAATCTTATT TCCTGCATAT TATACAGATA ACATAATGAA 4651 AGAAAAAATC TTTTTTTTTG TTCTTCAATG ATGATTTCAA CCATTCTTTT 4701 AAACATTGAT CAATTCCTGA GCAACAACCC CATACACACT GGTTTATATA 4751 CCGCCCCTTT TACAGTTGAA GAAAGAAATA GAAATAGAAA TAGCAAACAA 4801 AAGATATGAC AGTCAACACT AAGACCTATA GTGAGAGAGC AGAAACTCAT 4851 GCCTCACCAG TAGCACAGCG ATTATTTCGA TTAATGGAAC TGAAGAAAAC 4901 CAATTTATGT GCATCAATTG ACGTTGATAC CACTAAGGAG TTCCTCGAGT 4951 TAATTGATAA ATTAGGTCCT TATGTATGCT TAATCAAGAC TCATATTGAT 5001 ATAATCAATG ATTTTTCCTA TGAATCCACT ATTGAACCAT TATTAGAACT 5051 TTCACGTAAA CATCAATTTA TGATTTTTGA AGATAGAAAA TTTGCTGATA 5101 TTGGTAATAC CGTAAAGAAA CAATATATTG GTGGAGTTTA TAAAATTAGT 5151 AGTTGGGCAG ATATTACCAA TGCTCATGGT GTCACTGGGA ATGGAGTGGT 5201 TGAAGGATTA AAACAGGGAG CTAAAGAAAC CACCACCAAC CAAGAGCCAA 5251 GAGGGTTATT GATGTTAGCT GAATTATCAT CAGTGGGATC ATTAGCATAT 5301 GGAGAATATT CTCAAAAAAC TGTTGAAATT GCTAAATCCG ATAAGGAATT 5351 TGTTATTGGA TYTATTGCCC AACGTGATAT GGGTGGCCAA GAAGAAGGAT

FIG. 13 (CONTINUED 4).

5401 TTGATTGGCT TATTATGACA CCTGGAGTTG GATTAGATGA TAAAGGTGAT 5451 GGATTAGGAC AACAATATAG AACTGTTGAT GAAGTTGTTA GCACTGGAAC 5501 TGATATTATC ATTGTTGGTA GAGGATTGTT TGGTAAAGGA AGAGATCCAG 5551 ATATTGAAGG TAAAAGGTAT AGAAATGCTG GTTGGAATGC TTATTTGAAA 5601 AAGACTGGCC AATTATAAAT GTGAAGGGGG AGATTTTCAC TTTATTAGAT 5651 TTGTATATAT GTAGAATAAA TAAATAAATA AGTTAAATAA ATAATTAAAT 5701 AAGGGTGGTA ATTATTACTA TTTACAATCA AAGGTGGTCC TTCTAGCTGT 5751 AATCCGGGCA GCGCAACGGA ACATTCATCA GTGTAAAAAT GGAATCAATA 5801 AAGCCCTGCG CTCATGAGCC CGAAGTGGCG AGCCCGATCT TCCCCATCGG 5851 TGATGTCGGC GATATAGGCG CCAGCAACCG CACCTGTGGC GCCGCAGCGC 5901 GCAGGGTCAG CCTGAATACG CGTTTAATGA CCAGCACAGT CGTGATGGCA 5951 AGGTCAGAAT AGCCCAAGTC GGCCGAGGGG CCTGTACAGT GAGGGAAGAT 6001 CTGATATTGA CGAAGAGGAA CCAATGTAAC GTTACACTGA AGAAAACACA 6051 CAATAAACGG GAAGAAACGG TGTAAAAGTG TGAAAATAAT TTTTGAATAT 6101 CATTTCCCTT GGTTTAATTC CAAACGAAAC GTGTTTTTT TAGAGAATGG 6151 GAATTCTTAT TGGATGTCTA GATTGTTTGT TTACTCCAGA CTGTGCACAA 6201 AAACGTTTGG ATGGATGATC AGAAGATATT TTTAGGCTTA GCTCTAAATA 6251 TAAGAAATGA TGCTTGAAAA ACCAGACAGA AATTGAGTTT CAAAAATTGG 6301 TAATGTGAGG TATTAGTCAA CTAACCAAAT AACAATGCAA ACCGGTTGAT 6351 ACATTCATT TTGAAAATAA TGAAACTGGA ATTGGATGAC CAGCACACAA 6401 ACACATAAAG TAATTATGGG AATTAGAAGC GAACATAGAG GAGTACTTGG 6451 CCACGAACAG AATACAAGTG GGAACACTAT TTTCTCCATT GTTTTAGTTC 6501 TGTTTTTTG TCAGCCTAGT TTTGTGCTAT GTGTAAAAAA TATTGCCAAG 6551 AAAAAAAGCI TGTTTTGTGG CCAGIGTCCG AAAAAAATTT TGGGGAATCT 6601 TCGGATTAAT TTATGTTTTC A

FIG. 14

MITDEQLNTI ALTFGFASII LIIIYHAIST NVHKLEDETP SSSFTRTNTT

ETTVASKKKK 51 60

FIG. 15

MLKTLTQTLR LTGKAFPKVR PALIRTYAAF DRSKPHVNIG TIGHVDHGKT

TLTAAITKVL AEQGGANFLD YGSIDRAPEE RARGITISTA HVEYETKNRH 51 100

YAHVDCPGHA DYIKNMITGA AQMDGAIIVV AATDGQMPQT REHLLLARQV

GVQDLVVFVN KVDTIDDPEM LELVEMEMRE LLSTYGFDGD NTPVIMGSAL 151 200

MALEDKKPEI GKEAILKLLD AVDEHIPTPS RDLEQPFLLP VEDVFSISGR 201

GTVVTGRVER GVLKKGEEIE IVGGFDKPYK TTVTGIEMFK KELDSAMAGD

NCGVLLRGVK RDEIKRGMVL AKPGTATSHK KFLASLYILT SEEGGRSTPF 301

GEGYKPQCFF RTNDVTTTFS FPEGEGVDHS QMIMPGDNIE MVGELIKSCP
400

LEVNORFNLR EGGKTVGTGL ITRIIE 401 426

F1G.16.

MEVTQRTQSQ 1	TQPTQQSPTT	QTQTQSKEDQ	NRICQLICST	GQFGNYDLNI 50
NDKTIVQGKM 51	TWYFGRDPNS	DLQVASSSRI	SNKHFQIWLN	FNDKSLWIKD 100
TSTNGTHLNN 101	SRLVKGSNYL	LNQGDEIAVG	VGRDEDVVRF	VVVFGDKYNP 150
AKLPDSTNTI 151	KDEGIYKDFI	VKNETIGQGA	FATVKKAIER	STGESYAVKI 200
INRRKALNTG 201	GGSAMAGVDR	ELSILERLNH	PNIVALKAFY	EDMDNYYIVM 250
ELVPGGDLMD 251	FVAANGAIGE	DATQVITKQI	LEGIAYVHNL	GISHRDLKPD 300
NILIMQDDPI 301	LVKIŢDFGLA	KFSDNSTFMK	TFCGTLAYVA	PEVITGKYGS 350
SQMESQQKDN 351	YSSLVDIWSL	GCLVYVLLTS	HLPFNGKNQQ	QMFAKIKRGE 400
FHEAPLNSYD 401	ISEDGRDFLQ	CCLQVNPKLR	MTAAEALKHK	WLQDLYEEDS 450
VKSLSLSQSQ 451	SQQSRKIDNG	IHIESLSKID	EDVMLRPLDS	ERNRKSSKQQ 500
DFKVPKRVIP 501	LSQHPATPLP	MSQPKKRPYQ	IDPRTNKKVD	LEEPSTSKKV 550
KLSDSVVAED 551	YLKLEPLANS	LFQETINISK	SPFSFGRNDT	CDCEIDDDRL 600
SKLHCVITKE	NDSIWLLDKS	TNSCLVNNTS	VGKGNKVLLR	GGEILHLFFD 650
PLSSQHIGFK	VVLVDQSSGE	CXIVEVLKQ	TSEEMNIIPL	ISGLSSISS 699

		F/G. 17.		ما موسود موسود
MGTSTSEALK 1	NIKNKQRRQK	VFAEIKHEKN	KQRHKQRAER	AKEERENPEL 50
REERIAANIP 51	DTIDSKRIYD	ETIAAEVEGD	DEFQSYFTNL	LEEPKILLTT 100
SANAKKPAYE 101	FADMIMDFLP	NVTFIKRKKE	YTMQDMAKYC	SNRDFTALLV 150
INEDKKKVNG 151	ITLINLPEGP	TFYFSITSIV	DGKRIKGHGK	AGDYLPEIVL 200
NNFNSRLGKT 201	VGRLFQSIFP	HKPELQGRQV	ITLHNQRDYI	FFRRHRYIFR 250
NEEKVGLQE	GPQFTLKLRRM	QKGVRGDVVW	EHRPDMERDK	KKFYL 295

FIG. 18.

MNSEKIIEVI IAIFLPPVAV FMKCGATTPL WINLVLCIFI WFPAILHALY
1 50

VVLKD
51

29/53 F1G. 19.

MTLGFDKFIS KVSTHRRQSE PSILEIAATN SQNKSRRLSM DNGHCYVRES

1

50

TNNHHHLNTV VDNLRQRAGS FSFISHHHNH HQNSHDNYTV DPLTSNGARI

100

SRSRSKSV GHGEAISPAY FSKNKTKDLV KQETAHIISK KLLNMLQDLD

150

LQNPIALKTI SQGSESKFCK IYVSNTNNCI YLPAASSTSF TYEDDENGGV

200

IIAEDRNDEM PTAVNNNTLS MDSINHSETD FSDSPPPPDL FSKMKSFHSP 201

250

NYLTSKIDSE CPIPHTFAVI VELTKDSLII KDLHFQFQSL TTILWPTGDA 251

300

YNRTHAKEKF TIGNMEWRTS LSDADYYINS SNSNDVKSKN LGPEDLINRT 301

350

REYKLIDIEE PNNSSNSLSD DDMDINNITS PLSTSPTSSS TSTNSTSNSL

400

GSDSYKAGLY VFLLPILLPE HIPASIVSIN GSLAHTLSVE CNKYTDKLNR

450

KSKVSASYNL PMVRTPPNIG NSIADKPIYV NRIWNDAVHY IITFPRKYVT

500

LGCEHMINVK LSPMVXDVVI KRIKANVLER ITYVSKNLSR EYDYDSEDPY 501

550

CIHPVSKENK VRERVVSLYE LKIKAKQSSG GHLEAYKQEV MKCPENNLLF

600

SCYEVENDIN NGNGNGNGNG NYNVKQYNKD QPMIATPLDI NVSLPFLTIM 601

650

SDSLIMTSAI EEEGSDSPHT SRRGSAVSMT DNNTTPSNNN PLSPFLGAVE

30/53

FIG. 19 (CONTINUED).

651

700

TNGASINEIG DHTLFPDSNF RHIEIKHRLQ VTFRISKPDS DNKMHHYEVV

750

IDTPIVLLSS KCQEDSPPPY SSV 751

773

FIG. 20.

 ${\tt MGEGTPSLGKRHNKSHTLCNRCGRRSFHVQKKTCSSCGYPAAKMRSHNWALKAKRRRTTGTGRMAYLKHV\\ {\tt TRRFKNGFQTGVAKAQTPSA}$

FIG. 21.

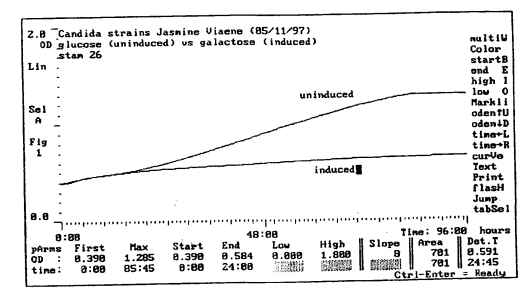
Project: Identification of novel essential genes in C. albicans

Strain no.: 26g

Freezer location: glycerol stocks box XXIII; C8

Growth curve(s) (Bioscreen):

Date: 05/11/1997



Plasmid/clone name*: 26g3

Freezer location: original stocks box

Identifier (gene name):

HTS screen:

Form generated by: Inge Loonen

FIG. 22.

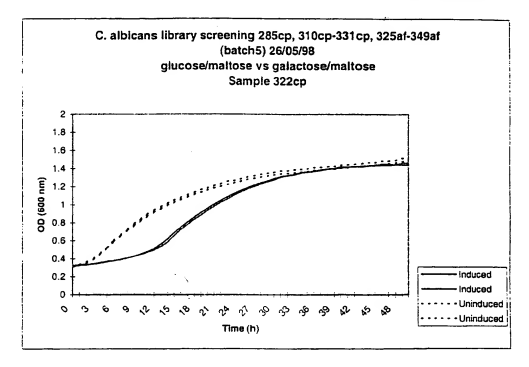
Project: Identification of novel essential genes in C. albicans

Strain no.: 322c_cp

Freezer location: glycerol stocks box XIV; D6

Growth curve(s) (Bioscreen):

Date: 26/05/1998



Plasmid/clone name*: 322c_cp Freezer location: original stocks box Identifier (gene name):

HTS screen:

Form generated by: Inge Loonen

F1G. 23.

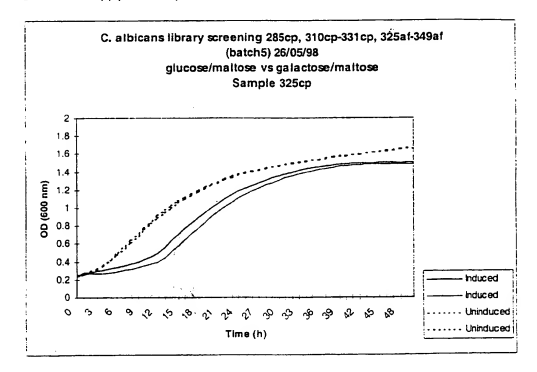
Project: Identification of novel essential genes in C. albicans

Strain no.: 325c_af

Freezer location: glycerol stocks box XIII; G4

Growth curve(s) (Bioscreen):

Date: 26/05/1998



Plasmid/clone name* : 325c_af
Freezer location : original stocks box

Identifier (gene name):

HTS screen:

Form generated by : Inge Loonen

FIG. 24.

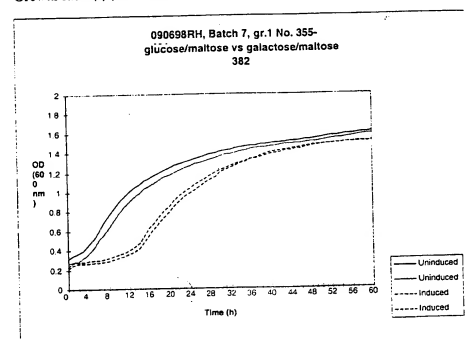
Project : Identification of novel essential genes in C. albicans

Strain no.: 382cp (FACS, batch 7, G1)

Freezer location: glycerol stocks box XVI; A2

Growth curve(s) (Bioscreen):

Date: 09/06/98



Plasmid/clone name*: 382cp (purified PCR product) Freezer location: original stocks box VIII; AAH8

Identifier (gene name): OST4

HTS screen:

Form generated by : Inge Loonen

35/53 F/G. 25.

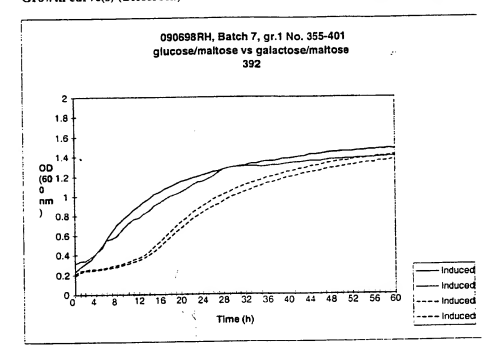
Project: Identification of novel essential genes in C. albicans

Strain no.: 392cp (FACS, batch 7, G1)

Freezer location: glycerol stocks box XVI; B3

Growth curve(s) (Bioscreen):

Date: 09/06/98



Plasmid/clone name*: 392cp (purified PCR product) Freezer location: original stocks box VIII; AAH2

Identifier (gene name): TUF1

HTS screen:

Form generated by: Inge Loonen

FIG. 26

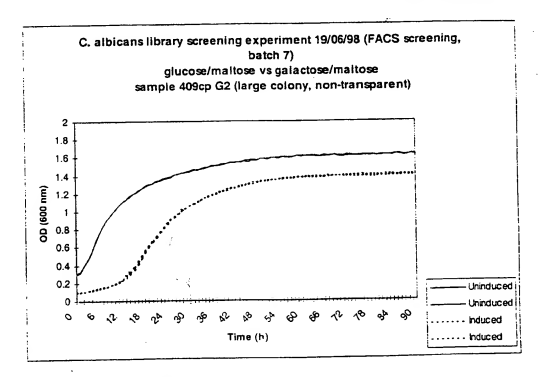
Project: Identification of novel essential genes in C. albicans

Strain no.: 409c_cp

Freezer location: glycerol stocks box XVI; C9

Growth curve(s) (Bioscreen):

Date: 19/06/1998



Plasmid/clone name*: 409c_cp Freezer location: original stocks box

Identifier (gene name):

HTS screen:

Form generated by: Inge Loonen

FIG. 27.

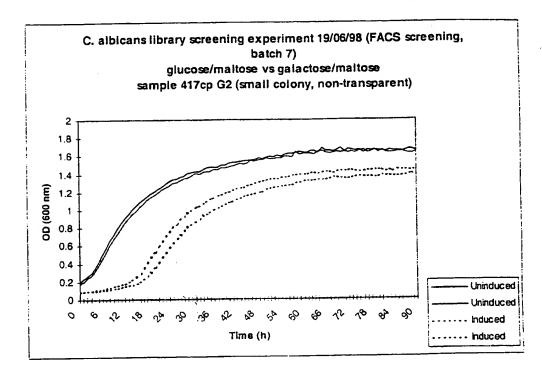
Project: Identification of novel essential genes in C. albicans

Strain no.: 417c_cpG2

Freezer location: glycerol stocks box XVI; D8

Growth curve(s) (Bioscreen):

Date: 19/06/1998



Plasmid/clone name*: 417c_cpG2L Freezer location: original stocks box

Identifier (gene name):

HTS screen:

Form generated by : Inge Loonen

FIG. 28.

Project : Identification of novel essential genes in C. albicans

Identifier (gene name) : 325caf

Disruptant strain:

Host strain:

Freezer location:

Disruption plasmid name*:

Freezer location:

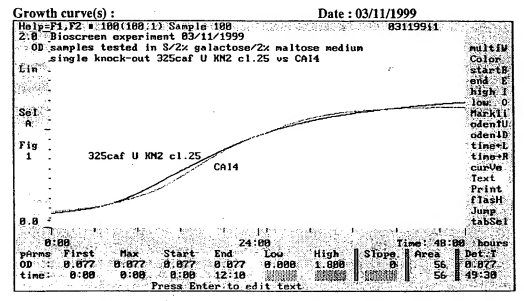
Knock-out (single/double):

Lab book ref.:

Southern results:

PCR results:

FIG. 28 (CONTINUED 1).



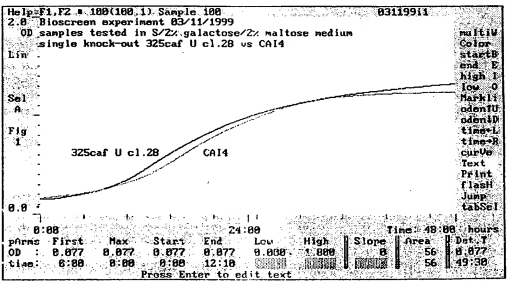
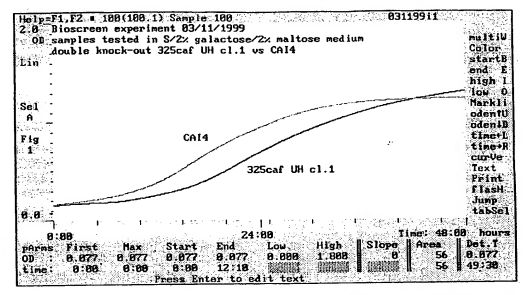
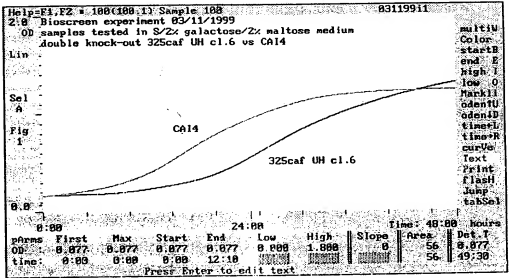


FIG. 28 (CONTINUED 2).





HTS screen:

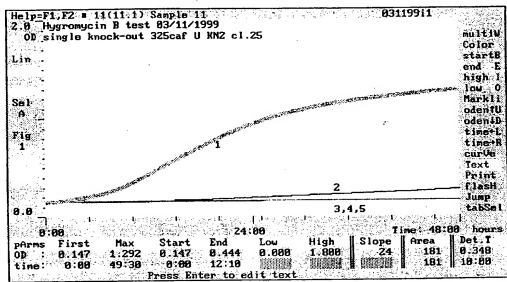
Bioscreen test of 325caf knock-out and WT growth in presence of hygromycin B

dilutions prepared

Stock solution of 53 mg/ml was prepared for hygromycin B. From this solution dilutions were prepared of: 4000µg/ml, 3000µg/ml, 2000µg/ml and 1000µg/ml

Growth curves for 325cafgK knock-out and WT in the presence of hygromycin B SUBSTITUTE SHEET (RULE 26)

FIG. 28 (CONTINUED 3).



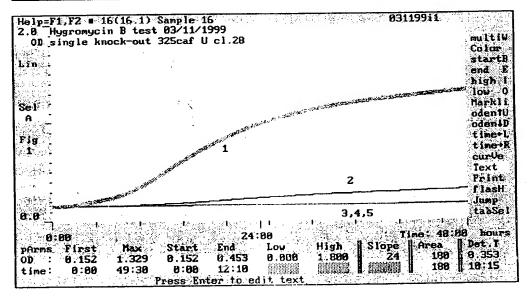
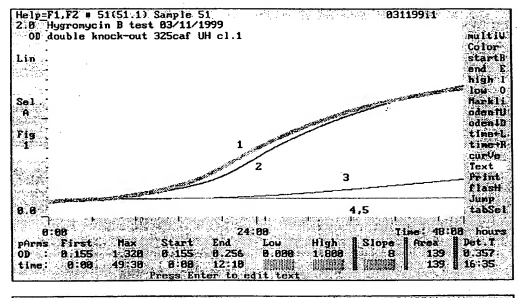


FIG. 28 (CONTINUED 4).



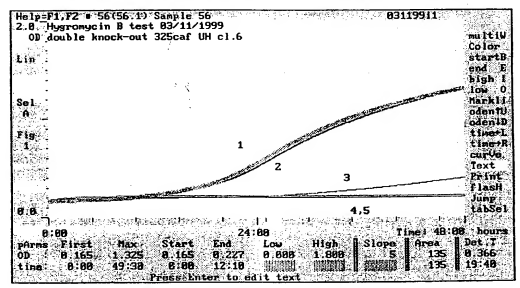
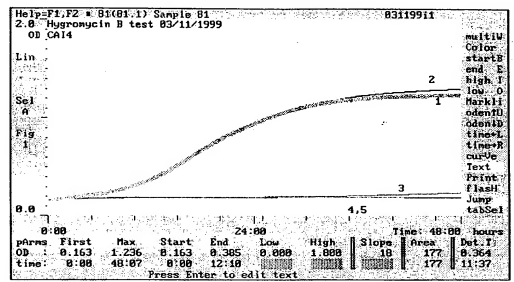


FIG. 28 (CONTINUED 5).



Legend:

- 1: S/2% gal/2% mal medium containing 0 μ g/ml Hygromycin B
- 2: S/2% gal/2% mal medium containing 1000 µg/ml Hygromycin B
- 3: S/2% gal/2% mal medium containing 2000 µg/ml Hygromycin B
- 4: S/2% gal/2% mal medium containing 3000 μg/ml Hygromycin B
- 5: S/2% gal/2% mal medium containing 4000 μg/ml Hygromycin B

Form generated by:

FIG. 29.

Project: Identification of novel essential genes in C. albicans

Identifier (gene name):

322c_cp

Disruptant strain:

322c_cp (in progress)

Host strain:

CAI4NG

Freezer location:

Knockout strain, box,

pos.

Disruption plasmid name*:

322c_cpURAcass.(inv)/pCR2.1(inv)

Freezer location:

single (in progress)

Knock-out (single/double): Lab book ref.:

Labbook 104 of Ronald de Hoogt

Southern results:

PCR results:

FIG. 30.

Project: Identification of novel essential genes in C. albicans

Identifier (gene name) : 417c_cpG2 Disruptant strain :

Host strain:

Freezer location:

Disruption plasmid name*:

Freezer location:

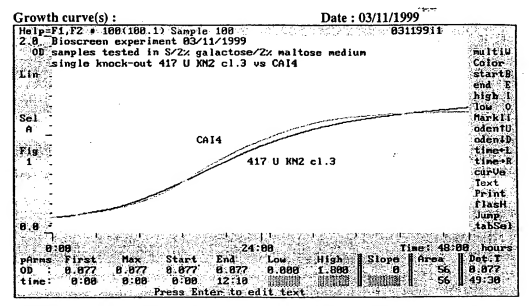
Knock-out (single/double):

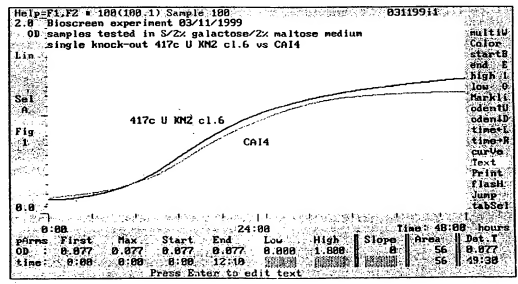
Lab book ref. :

Southern results:

PCR results:

FIG. 30 (CONTINUED 1).





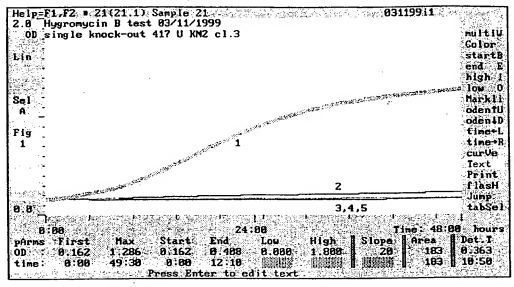
HTS screen:

Bioscreen test of 417c_cp knock-out and WT growth in presence of hygromycin B

dilutions prepared

Stock solution of 53 mg/ml was prepared for hygromycin B. From this solution dilutions were prepared of: 4000µg/ml, 3000µg/ml, 2000µg/ml and 1000µg/ml

FIG. 30 (CONTINUED 2).



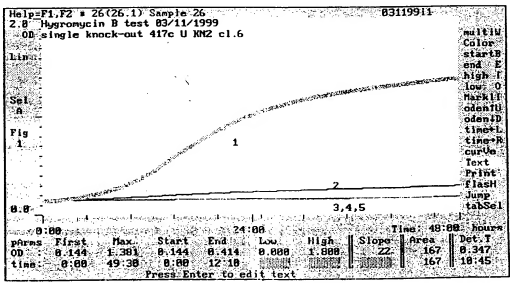
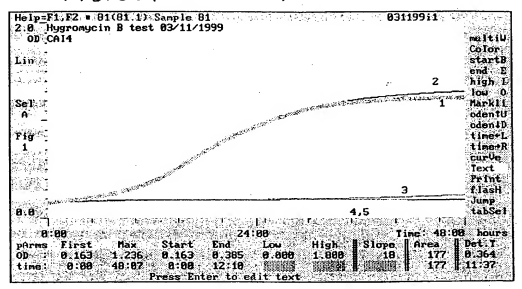


FIG. 30 (CONTINUED 3).



Legend:

- 1: S/2% gal/2% mal medium containing 0 µg/ml Hygromycin B
- 2: S/2% gal/2% mal medium containing 1000 µg/ml Hygromycin B
- 3: S/2% gal/2% mal medium containing 2000 µg/ml Hygromycin B
- 4: S/2% gal/2% mal medium containing 3000 µg/ml Hygromycin B
- 5: S/2% gal/2% mal medium containing 4000 µg/ml Hygromycin B

Form generated by:

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FIG. 31.

Project: Identification of novel essential genes in C. albicans

Identifier (gene name): TUF1

Disruptant strain: TUF1SAKO 7 Host strain: CAI4/NG

Freezer location: Strain collection Roland Contreras. YA132

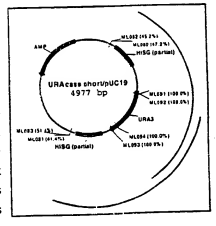
Disruption plasmid name*: Not applicable: short terminal homology (STH) PCR with overlapping fragments (split marker).

ML080	acataatcaagtgaatttacttacatcatttattgtggaaatttccgaatGTGCTGGAATTCGCCCTTTATG
ML081	tcacctatatatacctcttttttttattattcacagtgcacattctgtCCGGCTCGTATGTTGTG TGG
ML094	CCAGTGCTAACAACTTCATCAACAGTT
ML092	GCCTCACCAGTAGCACAACG

Uppercase sequences are segments that annual to the template DNA URAcass short/pUC19; the lowercase sequences are 50 nt uptstream (ML080), resp. downstream (ML081) of the target ORF.



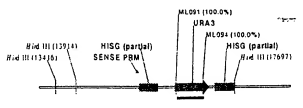
Amplification of STH fragments for TUF1 (ML080/ML094 \Rightarrow TUF1_STH-5'URA3 and ML081/ML092 \Rightarrow TUF1_STH-3'URA3). TUF1_STH-5'URA3 has a 50 bp terminal homology region upstream of the TUF1 ORF and a 3' incomplete URA3 marker; while TUF1_STH-3'URA3 has a 50 bp terminal homology region downstream of the TUF1 ORF and a 5' incomplete URA3 marker. In vivo, only an intact URAblast cassette can be formed when recombination occurs between the overlapping truncated URA3 sequences of the respective STH fragments.



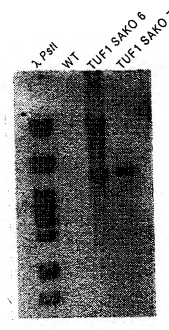
Southern results:

FIG. 31 (CONTINUED 1).

Presentation of disrupted allele



Fragment of TUF1 SAKO



HindIII digest URA3 probe Expected band: 3783 bp

PCR results:

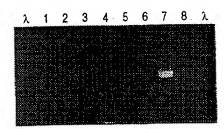
PCR analysis was performed using the primer combination ML091/ML093 (see figure), amplifying a URA3 fragment. Band of 755 bp points to correct homologous recombination of URA3 overlapping fragments. TUF1 SAKO 7 is clearly positive (SAKO stands for single allele knock out).



lane 1: λ Pst1 lane 2 to 9: potential TUF1 SAKCs 1 to 8 lane 10: Uracass short/pUC19 (positive control) lane 11: water lane 12: CAI4

51/53 FIG. 31(CONTINUED 2).

To check correct integration into the genome, PCR was performed with primer sets ML090/ML097. For TUF1 SAKO 7 a clear signal was obtained of the correct length of 1825 bp.



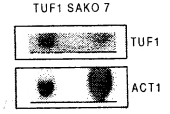
λ: λ Pstl 1 to 8: potential TUF1 SAKOs 1 to 8

Date: 03/11/1999

Northern analysis:

Growth curve(s):

88% inhibition



Help=F1,F2 = 188(188.1) Sample 188

2.8 Bioscreen experiment 83/11/1999

OB samples tested in S/2% galactose/2% maltose medium single knock-out TUF1 cl.7 vs CAl4 03119911 multiU Color startB end high low Sel Markli odentU A odenID time+L CAI4 time R curve TUF1 cl.7 Text Print f lasH Jump. tabSel 0.6 Time: 48:88 | hours 24:88 0:00 Det.T 0.077 High pArms OD : Start I.ou First Max.

0.077

8:80

8.877

0:00

0.077

9:00

OD

time:

0.877

12:10

868.8

1.800

56

FIG. 31 (CONTINUED 3).

Bioscreen test of TUF1 knock-out clone 7 and WT growth in presence of hygromycin B

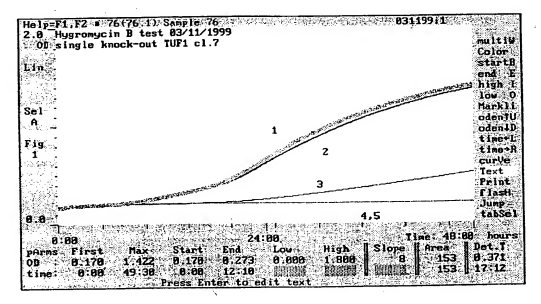
dilutions prepared

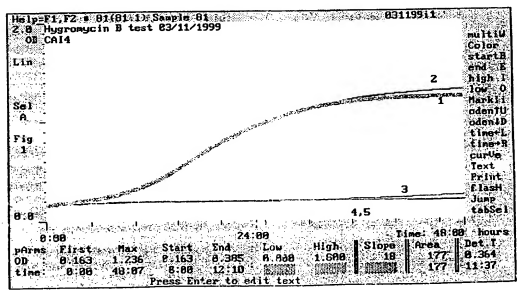
Stock solution of 53 mg/ml was prepared for hygromycin B.

From this solution dilutions were prepared of:

 $4000\mu g/ml$, $3000\mu g/ml$, $2000\mu g/ml$ and $1000\mu g/ml$

Growth curves for TUF1 knock-out and WT in the presence of hygromycin B





Legend:

FIG. 31 (CONTINUED 4).

- 1: S/2% gal/2% mal medium containing 0 μ g/ml Hygromycin B
- 2: S/2% gal/2% mal medium containing 1000 μ g/ml Hygromycin B
- 3: S/2% gal/2% mal medium containing 2000 μ g/ml Hygromycin B
- 4: S/2% gal/2% mal medium containing 3000 μ g/ml Hygromycin B
- 5: S/2% gal/2% mal medium containing 4000 µg/ml Hygromycin B

SEQUENCE LISTING

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<110> Janssen Pharmaceutica N. V.
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tcataatata tcatgccata tctactaatg tacataaatt agaagatgaa accccatcat 180
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PCT/EP99/09833 WO 00/34481

caccgttacc aatgtcacaa ccgaaaaaga ggccgtatca aatagaccct agaacaaaca 1680 aaaaagtcga tttggaagaa cctctgacaa gcaagaaagt caagctaagt gattccgttg 1740 ttgcggaaga ctacttgaag ttggggccac ttgcaaattc gttattccaa gaaacaataa 1800 atatttcaaa gtccccgttt tctttcggaa gaaatgacac ttgtgattgc gagatagacg 1860 acgacagact atccaaactt cattgtgtca ttaccaaaga aaacgactct atatggttat 1920 tggataagag tactaactcg tgcttggtca acaatactag tgttggaaaa ggcaacaaag 1980 ttttgcttag aggagggag atattacatc tcttctttga cccattgtca ctgcaacata 2040 taggtttcaa agtagtcctt gttgatcaac tgtctggtga acataagagt caagtggagg 2100 ttttgaaaca aacctcagaa gaaatgaata ttattccact tatttctggt ttaagtagta 2160 taagttoata gatttagoat atatacaago atttootata gaaacaaagg ttoattaatt 2220 tagttattta cetecatgea attacattta ettettette caagggegaa ttetgeagat 2280 2283

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aataatttca attcaagatt gggtaaaact gtgggaagac tatttcaaag tattttccct 660
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ggtccgcagt ttacattaaa gctaagaaga atgcaaaagg gagtacgtgg tgatgttgtt 840
tgggaacaca gaccagatat ggaaagagat aagaagaagt tttatttata agc<del>ggg</del>tgta 900
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 gctgtgttta tgaaatgtgg tgccactacc ccattatgga ttaacttggt attatgtatc 180
 tttatttggt tecetgetat ettacatgee ttataegttg tgttgaaaga ttaaacaaac 240
 accagagatt tactgcttga tgaattgatt actccaaaga gttgtgacta gttcccagtg 300
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 ctattttgca gagttttcaa aatttatcca aaacatgtta gtcattaaac catattatta 420
 gtttggtatt ttatttttt gtatttatca attggaatat atatctatac atgaatttat 540
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  gataatggtc attgttatgt tcgtgaatca actaataatc atcatcattt aaataccgtc 180
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  catcagaata gtcacgataa ttatactgtc gatcccctta catcaaacgg agcacgaatt 300
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  ttttccaaga ataaaaccaa agatttagtg aaacaggaaa cagcacatat cattctgaag 420
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ccatagttca gaaaataaaa ttgaaaaatt taaaaaaaa cgcaaatatca ttcattttt 180
ttgtttttt gacaataata ttaatatgta gttaccaatg tttttagatt ttatatgttt 240
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<210> 9 <211> 119 <212> DNA <213> Candida albicans

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<211> 60

<212> PRT

<213> Candida albicans

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His Lys Leu Glu Asp Glu Thr Pro Ser Ser Ser Phe Thr Arg Thr Asn 40 35

Thr Thr Glu Thr Thr Val Ala Ser Lys Lys Lys 55

<210> 11

<211> 426

<212> PRT

<213> Candida albicans

<400> 11

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Pro Lys Val Arg Pro Ala Leu Ile Arg Thr Tyr Ala Ala Phe Asp Arg 25 20

Ser Lys Pro His Val Asn Ile Gly Thr Ile Gly His Val Asp His Gly 40 35

Lys Thr Thr Leu Thr Ala Ala Ile Thr Lys Val Leu Ala Glu Gln Gly 55

Gly Ala Asn Phe Leu Asp Tyr Gly Ser Ile Asp Arg Ala Pro Glu Glu 75 70

Arg Ala Arg Gly Ile Thr Ile Ser Thr Ala His Val Glu Tyr Glu Thr 90 85

Lys Asn Arg His Tyr Ala His Val Asp Cys Pro Gly His Ala Asp Tyr 105 100

Ile Lys Asn Met Ile Thr Giy Ala Ala Gin Met Asp Gly Ala Ile Ile 125 12C

Val Val Ala Ala Thr Asp Gly Gln Met Pro Gln Thr Arg Glu His Leu 130 135 140

- Leu Leu Ala Arg Gln Val Gly Val Gln Asp Leu Val Val Phe Val Asn 145 150 150 155 160
- Lys Val Asp Thr Ile Asp Asp Pro Glu Met Leu Glu Leu Val Glu Met 165 170 175
- Glu Met Arg Glu Leu Leu Ser Thr Tyr Gly Phe Asp Gly Asp Asn Thr 180 185 190
- Pro Val Ile Met Gly Ser Ala Leu Met Ala Leu Glu Asp Lys Lys Pro 195 200 205
- Glu Ile Gly Lys Glu Ala Ile Leu Lys Leu Leu Asp Ala Val Asp Glu 210 215 220
- His Ile Pro Thr Pro Ser Arg Asp Leu Glu Gln Pro Phe Leu Leu Pro 225 230 230 235
- Val Glu Asp Val Phe Ser Ile Ser Gly Arg Gly Thr Val Val Thr Gly 245 250
- Arg Val Glu Arg Gly Val Leu Lys Lys Gly Glu Glu Ile Glu Ile Val 260 265 270
- Gly Gly Phe Asp Lys Pro Tyr Lys Thr Thr Val Thr Gly Ile Glu Met 275 280 285
- Phe Lys Lys Glu Leu Asp Ser Ala Met Ala Gly Asp Asn Cys Gly Val 290 295 300
- Leu Leu Arg Gly Val Lys Arg Asp Glu Ile Lys Arg Gly Met Val Leu 305 310 315 320
- Ala Lys Pro Gly Thr Ala Thr Ser His Lys Lys Phe Leu Ala Ser Leu 325
- Tyr Ile Leu Thr Ser Glu Glu Gly Gly Arg Ser Thr Pro Phe Gly Glu 340
- Gly Tyr Lys Pro Gln Cys Phe Phe Arg Thr Ash Asp Val Thr Thr Thr 355
- Phe Ser Phe Pro Glu Gly Glu Gly Val Asp Ris Ser Gln Met Ile Met 370 375 380

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Pro Gly Asp Asn Ile Glu Met Val Gly Glu Leu Ile Lys Ser Cys Pro 395 390 385

Leu Glu Val Asn Gln Arg Phe Asn Leu Arg Glu Gly Gly Lys Thr Val 410 405

بمييود Gly Thr Gly Leu Ile Thr Arg Ile Ile Glu 425 420

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<211> 699

<212> PRT

<213> Candida albicans

<400> 12

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Ile Cys Gln Leu Ile Cys Ser Thr Gly Gln Phe Gly Asn Tyr Asp Leu 40 35

Asn Ile Asn Asp Lys Thr Ile Val Gln Gly Lys Met Thr Trp Tyr Phe 55

Gly Arg Asp Pro Asn Ser Asp Leu Gln Val Ala Ser Ser Ser Arg Ile 70 .

Ser Asn Lys His Phe Gln Ile Trp Leu Asn Phe Asn Asp Lys Ser Leu 90 85

Trp Ile Lys Asp Thr Ser Thr Asn Gly Thr His Leu Asn Asn Ser Arg 105 100

Leu Val Lys Gly Ser Asn Tyr Leu Leu Asn Gln Gly Asp Glu Ile Ala 120 115

Val Gly Val Gly Arg Asp Glu Asp Val Val Arg Phe Val Val Phe 135 130

Gly Asp Lys Tyr Asn Pro Ala Lys Leu Pro Asp Ser Thr Asn Thr Ile 155 150 145

Lys Asp Glu Gly Ile Tyr Lys Asp Phe Ile Val Lys Asn Glu Thr Ile 170 165

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Gly	Gln	Gly	Ala 180	Phe	Ala	Thr	Val	Lys 185	Lys	Ala	Ile	Glu	Arg 190	Ser	Thr	
Gly	Glu	Ser 195	Tyr	Ala	Val	Lys	Ile 200	Ile	Asn	Arg	Arg	Lys 205	Ala	Leu		
Thr	Gly 210	Gly	Gly	Ser	Ala	Met 215	Ala	Gly	Val	Asp	Arg 220	Glu	Leu	Ser	Ile	!
Leu 225	Glu	Arg	Leu	Asn	His 230	Pro	Asn	Ile	Val	Ala 235	Leu	Lys	Ala	Phe	Tyr 240	:)
Glu	Asp	Met	Asp	Asn 245		Tyr	Ile	Val	Met 250	Glu	Leu	Val	Pro	Gly 255	Gly	Y
Asp	Lev	ı Met	260		e Val	Ala	Ala	Asr 265	Gly	, Ala	Ile	: Gly	7 Glu 270	. Asp	Al	a
Thr	Gl:	n Val		e Thi	: Lys	Glr	11e		ı Glı	ı Gly	/ Ile	285	a Tyr	: Val	. Hi	s
	29	0				29	5			s Pro	301	U				
30	5				31	0				e Th:	5				J.	
Ly	s Ph	e Se	r As	p As 32		r Th	r Ph	e Me	t Ly 33	rs Th	r Ph	е Су	s Gl	y Th 33	r Le 5	eu
Al	a Ty	r Va	1 Al 34		:o Gl	u Va	11 11	e Th	nr G] 15	Ly Ly	s Ty	r Gl	y Se. 35	r Se iO	r G	ln
M∈	et G		er Gl	ln G	ln Ly	/s As	sp As	sn Ty 60	yr S	er Se	er Le	eu Va 30	al As 65	;p Il	.e T	rp
Se		eu G 70	ly C	ys L	eu Va		yr V. 75	al L	eu L	eu Th	nr Se	er H: 80	is Le	eu Pr	o P	he
	sn G 85	ly L	ys A	sn G		ln G 90	in M	et F	he A	la L;	ys I 95	le L	ys A	rg G	ly C	31u 400
P	he H	is G	lu A		ro È 05	eu A	sn S	er T	'yr P	sp I	le S	er G	lu A	sp G 4	ly 1 15	Arg
A	sp E	Phe I	Leu C	iln (ys C	ys <u>I</u>	_eu (Sln V	al A	Asr. P	ro ï	.ys I	Leu A	Arg M	let	Thr

Ala Ala Glu Ala Leu Lys His Lys Trp Leu Gln Asp Leu Tyr Glu Glu Asp Ser Val Lys Ser Leu Ser Leu Ser Gln Ser Gln Ser Gln Ser Arg Lys Ile Asp Asn Gly Ile His Ile Glu Ser Leu Ser Lys Ile Asp Glu Asp Val Met Leu Arg Pro Leu Asp Ser Glu Arg Asn Arg Lys Ser Ser Lys Gln Gln Asp Phe Lys Val Pro Lys Arg Val Ile Pro Leu Ser 510 -Gln His Pro Ala Thr Pro Leu Pro Met Ser Gln Pro Lys Lys Arg Pro Tyr Gln Ile Asp Pro Arg Thr Asn Lys Lys Val Asp Leu Glu Glu Pro Ser Thr Ser Lys Lys Val Lys Leu Ser Asp Ser Val Val Ala Glu Asp Tyr Leu Lys Leu Gly Pro Leu Ala Asn Ser Leu Phe Gln Glu Thr Ile Asn Ile Ser Lys Ser Pro Phe Ser Phe Gly Arg Asn Asp Thr Cys Asp Cys Glu Ile Asp Asp Asp Arg Leu Ser Lys Leu His Cys Val Ile Thr Lys Glu Asn Asp Ser Ile Trp Leu Leu Asp Lys Ser Thr Asn Ser Cys Leu Val Asn Asn Thr Ser Val Gly Lys Gly Asn Lys Val Leu Leu Arg Gly Gly Glu Ile Leu His Leu Phe Phe Asp Pro Leu Ser Ser Gln His Ile Gly Phe Lys Val Val Leu Val Asp Gln Ser Ser Gly Glu His Lys Ser Gln Val Glu Val Leu Lys Gln Thr Ser Glu Glu Met Asn Ile Ile

Pro Leu Ile Ser Gly Leu Ser Ser Ile Ser Ser 690 695

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<211> 295

<212> PRT

<213> Candida albicans

<400> 13

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Arg His Lys Gln Arg Ala Glu Arg Ala Lys Glu Glu Arg Glu Asn Pro 35 40 45

Glu Leu Arg Glu Glu Arg Ile Ala Ala Asn Ile Pro Asp Thr Ile Asp 50 55 60

Ser Lys Arg Ile Tyr Asp Glu Thr Ile Ala Ala Glu Val Glu Gly Asp
65 70 75 80

Asp Glu Phe Gln Ser Tyr Phe Thr Asn Leu Leu Glu Glu Pro Lys Ile 85 90 95

Leu Leu Thr Thr Ser Ala Asm Ala Lys Lys Pro Ala Tyr Glu Phe Ala 100 105 110

Asp Met Ile Met Asp Phe Leu Pro Asn Val Thr Phe Ile Lys Arg Lys 115 120 125

Lys Glu Tyr Thr Met Gln Asp Met Ala Lys Tyr Cys Ser Asn Arg Asp 130 135 140

Phe Thr Ala Leu Leu Val Ile Asn Glu Asp Lys Lys Lys Val Asn Gly 145 150 150

Ile Thr Leu Ile Asn Leu Prc Glu Gly Pro Thr Phe Tyr Phe Ser Ile 165 170 175

Thr Ser Ile Val Asp Gly Lys Arg Ile Lys Gly His Gly Lys Ala Gly 180 185 190

Asp Tyr Leu Pro Glu Tie Val Leu Asn Asn Phe Asn Ser Arg Leu Gly

195 200 205

Lys Thr Val Gly Arg Leu Phe Gln Ser Ile Phe Pro His Lys Pro Glu 210 215 220

Leu Gln Gly Arg Gln Val Ile Thr Leu His Asn Gln Arg Asp Tyr Ile
225 230 235 240

Phe Phe Arg Arg His Arg Tyr Ile Phe Arg Asn Glu Glu Lys Val Gly 245 250 255

Leu Gln Glu Gly Pro Gln Phe Thr Leu Lys Leu Arg Arg Met Gln Lys 260 265 270

Gly Val Arg Gly Asp Val Val Trp Glu His Arg Pro Asp Met Glu Arg 275 280 285

Asp Lys Lys Lys Phe Tyr Leu 290 295

<210> 14

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<212> PRT

<213> Candida albicans

<400> 14

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Leu Tyr Val Val Leu Lys Asp 50 - 55

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<211> 773

<212> PRT

<213> Candida albicans

<400> 15

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1 5 10 15

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Asn	Lys	Ser 35	Arg	Arg	Leu	Ser	Met 40	Asp	Asn	Gly	His	Cys 45	Tyr	Val	
Glu	Ser 50	Thr	Asn	Asn	His	His 55	His	Leu	Asn	Thr	Val 60	Val	Asp	Asn	Leu
Arg 65	Gln	Arg	Ala	Gly	Ser 70	Phe	Ser	Phe	Ile	Ser 75	His	His	His	Asn	His 80
His	Gln	Asn	Ser	His 85	Asp	Asn	Tyr	Thr	Val 90	Asp	Pro	Leu	Thr	Ser 95	Asn
Gly	Ala	Arg	Ile 100	Ser	Arg	Ser	Arg	Ser 105	Arg	Ser	Lys	Ser	Val 110	Gly	His
Gly	Glu	Ala 115	Ile	Ser	Pro	Ala	Tyr 120	Phe	Ser	Lys	Asn	Lys 125	Thr	Lys	Asp
Leu	Val 130	Lys	Gln	Glu	Thr	Ala 135	His	Ile	Ile	Ser	Lys 140	Lys	Leu	Leu	Asn
Met 145	Leu	Gln	Asp	Leu	Asp 150	Leu	Gln	Asn	Pro	Ile 155	Ala	Leu	Lys	Thr	Ile 160
Ser	Gln	Gly	Ser	Glu 165	Ser	Lys	Phe	Cys	Lys 170	Ile	Tyr	Val	Ser	Asn 175	Thr
Asn	Asn	Cys	Ile 180	Tyr	Leu	Pro	Ala	Ala 185	Ser	Ser	Thr	Ser	Phe 190	Thr	Tyr
Glu	Asp	Asp 195		Asn	Gly	Gly	Val 200	Ile	Ile	Ala	Glu	Asp 205	Arg	Asn	Asp
Glu	Met. 210		Thr	Ala	Val	Asn 215		Asn	Thr	Leu	Ser 220	Met	Asp	Ser	Ile
Asn 225		Ser	Glu	Thr	Asp 230		Ser	Asp	Ser	Pro 235		Pro	Pro	Asp	Leu 240
Phe	Ser	Lys	Met	Lys 245		Phe	His	Ser	Pro 250		Tyr	Leu	Thr	Ser 255	Lys
Ile	Asp	Ser	Glu 260		Pro	Ile	Pro	His 265		: Phe	Ala	Val	11e		Glu

Leu Thr Lys Asp Ser Leu Ile Ile Lys Asp Leu His Phe Gln Phe Gln

Ser Leu Thr Thr Ile Leu Trp Pro Thr Gly Asp Ala Tyr Asn Arg Thr

280

275

	290					295					300			**	
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Leu	Ser	Asp	Ala	Asp 325	Tyr	Tyr	Ile	Asn	Ser 330	Ser	Asn	Ser	Asn	Asp 335	Val
Lys	Ser	Lys	Asn 340	Leu	Gly	Pro	Glu	Asp 345	Leu	Ile	Asn	Arg	Thr 350	Arg	Glu
Tyr	Lys	Leu 355	Ile	Asp	Ile	Glu	Glu 360	Pro	Asn	Asn	Ser	Ser 365	Asn	Ser	Leu
Ser	Asp 370	Asp	Asp	Met	Asp	Ile 375	Asn	Asn	Ile	Thr	Ser 380	Pro	Leu	Ser	Thr
Ser 385	Pro	Thr	Ser	Ser	Ser 390	Thr	Ser	Thr	Asn	Ser 395	Thr	Ser	Asn	Ser	Leu 400
Gly	Ser	Asp	Ser	Tyr 405	Lys	Ala	Gly	Leu	Tyr 410	Val	Phe	Leu	Leu	Pro 415	Ile
Leu	Leu	Pro	Glu 420	His	Ile	Pro	Ala	Ser 425	Ile	Val	Ser	Ile	Asn 430	Gly	Ser
Leu	Ala	His 435	Thr	Leu	Ser	Val	Glu 440	Cys	Asn	Lys	Tyr	Thr 445	Asp	Lys	Leu
Asn	Arg 450	Lys	Ser	Lys	Val	Ser 455	Alā	Ser	Tyr	Asn	Leu 460	Pro	Met	Val	Arg
Thr 465	Pro	Pro	Asn	Ile	Gly 470	Asn	Ser	Ile	Ala	Asp 475	Lys	Pro	Ile	Tyr	Val 480
Asn	Arg	Ile	Trp	Asn 485	Asp	Ala	Val	His	Tyr 490		Ile	Thr	Phe	Pro 495	Arg
Lys	Tyr	Val	Thr 500	Leu	Gľy	Cys	Glu	His 505		Ile	Asn	Val	Lys 510	Leu	Ser
Pro	Met	Val 515		Asp	Val	Val	Ile 520		Arg	Ile	Lys	Phe 525		Val	Leu
							14								

Glu Arg Ile Thr Tyr Val Ser Lys Asn Leu Ser Arg Glu Tyr Asp Tyr Asp Ser Glu Asp Pro Tyr Cys Ile His Pro Val Ser Lys Glu Asn Lys Val Arg Glu Arg Val Val Ser Leu Tyr Glu Leu Lys Thr Lys Ala Lys Gln Ser Ser Gly Gly His Leu Glu Ala Tyr Lys Gln Glu Val Met Lys Cys Pro Glu Asn Asn Leu Leu Phe Ser Cys Tyr Glu Val Glu Asn Asp Asn Asn Asn Gly Asn Gly Asn Gly Asn Gly Asn Gly Asn Lys Asn Val Lys Gln Lys Asn Lys Asp Gln Pro Met Ile Ala Thr Pro Leu Asp Ile Asn Val Ser Leu Pro Phe Leu Thr Thr Met Ser Asp Ser Leu Ile Met Thr Ser Ala Ile Glu Glu Glu Gly Ser Asp Ser Pro His Thr Ser Arg Arg Gly Ser Ala Val Ser Met Thr Asp Asn Asn Thr Thr Pro Ser Asn Asn Asn Pro Leu Ser Pro Phe Leu Gly Ala Val Glu Thr Asn Gly Ala Ser Ile Asn Glu Ile Gly Asp His Thr Leu Phe Pro Asp Ser Asn Phe Arg His Ile Glu Ile Lys His Arg Leu Gln Val Thr Phe Arg Ile Ser Lys Pro Asp Ser Asp Asn Lys Met His His Tyr Glu Val Val Ile Asp Thr Pro Ile Val Leu Leu Ser Ser Lys Cys Gln Glu Asp Ser Pro Pro Pro Tyr Ser Ser Val

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<213> Candida albicans

<400> 16

Met Gly Glu Gly Thr Fro Ser Leu Gly Lys Arg His Asn Lys Ser His 1 5 10 15

Thr Leu Cys Asn Arg Cys Gly Arg Arg Ser Phe His Val Gln Lys Lys 20 25 30

Thr Cys Ser Ser Cys Gly Tyr Pro Ala Ala Lys Met Arg Ser His Asn 35 40 45

Trp Ala Leu Lys Ala Lys Arg Arg Arg Thr Thr Gly Thr Gly Arg Met 50 55 60

Ala Tyr Leu Lys His Val Thr Arg Arg Phe Lys Asn Gly Phe Gln Thr 65 75 80

Gly Val Ala Lys Ala Gln Thr Pro Ser Ala 85 90

Comprehendance